# nuXmv 1.0 User Manual 

Marco Bozzano, Roberto Cavada, Alessandro Cimatti, Michele Dorigatti, Alberto Griggio, Alessandro Mariotti, Andrea Micheli, Sergio Mover, Marco Roveri, Stefano Tonetta

FBK - Via Sommarive 18, 38055 Povo (Trento) - Italy

Email: nuxmv@list.fbk.eu


This document is part of the distribution package of the NUXMV model checker.
For any additional request for information please send an e-mail to:

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$\underset{\substack{\text { Fondozione } \\ \text { BRUNO KESSER }}}{ }$

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## Chapter 1

## Introduction

nUXMV inherits, and thus provides to the user, all the functionalities of NUSMV [CCG ${ }^{+} 02$ ]. In this section we revise all the new features distinguishing them in those for the analysis of finite-state domains, those for the analysis of infinite-state domains, and other generic features.

### 1.1 Analysis of finite-state domains

nUXMV complements the NUSMV language with the aiger 1.9.4 [BHW11] format. aiger 1.9.4 is the language adopted in the hardware model checking competition. Once the aiger 1.9.4 file is read, the internal data structures of NUXMV are populated, and it is possible to verify the properties (if any) with any of the available verification algorithms, or specify new properties interactively "playing" with the design.
nUXmV implements a vast portfolio of algorithms for invariant checking. We extended MiniSat [ES03] to build a resolution proof. This enables for the extraction of interpolants [McM04], and opens for the implementation of interpolation based algorithms. We currently provide an implementation for the McMilllan approach [McM03] and for the interpolation sequence approach [VG09]. Interpolation based algorithms are complemented with k-induction algorithms [SSS00] and a family of algorithms based on IC3 [Bra11, HBS13, VGS12]. The IC3 algorithm using abstraction refinement [VGS12] comes in two variant depending on the approach to refinement: the original one based on IC3, and a new variant based on BMC. All these techniques, benefit from the use of temporal decomposition [CMBK09] and from the techniques to discover equivalences to simplify the problem.

Still related to the verification of invariants, we also improve the BDD based invariant checking algorithms by allowing the user to specify hints in the spirit of guided reachability [TCP08]. The hints are specified using a restricted fragment of the PSL SERE [EF06]. The hints can also be used to compute the full set of the reachable states.

For LTL SAT based model checking, we complement the BMC based algorithms of NuSMV [BCCZ99b, $\mathrm{BHJ}^{+}$06] with k-liveness [CS12] integrated within an IC3 framework. K-liveness is based on counting and bounding the number of times a fairness constraint can become true. This is used in conjunction with the construction of a monitor for LTL properties, for which we use the LTL2SMV [CGH97b] as provided by NUSMV.

### 1.2 Analysis of infinite-state domains

In order to allow the user to specify infinite-state systems, we extend the language of NUSMV with two new data types, namely Reals and unbounded Integers. This, for instance, enables to specify domains with infinite data types (e.g. the example in Fig. 1.1).

To analyze such kind of designs, we integrate in nuXmv several new verification algorithms based on Satisfiability Modulo Theory (SMT) [BSST09] and on abstraction, or combination of abstraction with other techniques.

We lift Simple Bounded Model Checking (SBMC) $\left[\mathrm{BHJ}^{+} 06\right]$ from the pure Boolean case to the SMT case. The encoding is the same as that of SBMC, but instead of using a SAT solver we use an SMT solver. The SBMC SMT based approach for LTL verification is complemented with k-liveness combined with IC3 extended

```
1 MODULE main
2 IVAR
3 d : Real;
4 VAR state:{s0, s1}
state :{s0, s1}
    ASSIGN
        init(state) := s0;
            next(state) := case
                state = s0 & res >= 0.10: s1;
                l
    esac;
    next(t):= case
        state = s0 & res < 0.10 : res + d;
        state = s1 & res < 0.20: res + d;
        TRUE
    INIT
    res >= 0.0
    TRANS
    (state = sO }->(d>=0 & d<= 0.01)) &
    (state =s1 }>(d>=0&d<=0.02)
INVARSPEC res <= 0.3
```

Figure 1．1：Example of the nuXmv language．
to the infinite－state case［CGMT14b］．This approach relies on recent results on applying an IC3－based approach to the verification of infinite－state systems［CG12］．We remark that，although these approaches are in general incomplete，if a lazo－shaped counterexample exists，it is guaranteed to be eventually found．Moreover，for certain designs，these approaches are able to conclude that the property hold．

As far as invariant checking is concerned，we lift the pure Boolean approaches like BMC，k－induction，inter－ polation，and IC3，to the case of verification of infinite－state systems．Intuitively，we use an SMT solver in place of a SAT solver．For the infinite case，similar to the finite case，we provide an SMT based implementation for McMilllan approach［McM03］；for the interpolation sequence approach［VG09］；for k－induction［SSS00］；and for a family of algorithms based on IC3［CG12，CGMT14a］．

NUXMV also implements several approaches based on abstraction refinement［CGJ ${ }^{+} 03$ ］．We provide new algorithms combining abstraction with BMC and k －induction［Ton09］．The algorithms do not rely on quantifier elimination techniques to compute the abstraction，but encode the model checking problem over the abstract state space into an SMT problem．The advantage，is that they avoid the possible bottleneck of abstraction computation． The very same approach has been recently lifted and tightly integrated within the IC3 framework［CGMT14a］， with very good results．All these techniques complement the＂classical＂counterexample guided（predicate）ab－ straction refinement（CEGAR）［CGJ $\left.{ }^{+} 03\right]$ ，also implemented in NUXmV．The CEGAR approach requires the computation of a quantifier－free formula that is equivalent to the abstract transition relation w．r．t．a given set of predicates．This，in turn，requires the solving of an ALLSAT problem［LNO06］．For this step，nUXmv imple－ ments different techniques：the combination of BDD and SMT［CCF $\left.{ }^{+} 07, \mathrm{CFG}^{+} 10\right]$ ，where BDDs are used as compact Boolean model enumerator within an ALLSMT approach；a technique that exploits the structure of the system under verification，by partitioning the abstraction problem into the combination of several smaller abstrac－ tion problems［CDJR09］．For the refinement step to discard the spurious counterexample，nUXmV implements three approaches based on the analysis of the unsatisfiable core，on the analysis of the interpolants，and on the weakest preconditions．

## 1．3 Miscellaneous functionalities

nUXMV provides novel functionalities that aim at facilitating the modeling and the understanding of complex designs．For instance，it allows for the generation of an explicit state representation（subject to the projection over a set of user specified predicates）in XMI format of the design under verification．The generated XMI can be visualized in any UML based viewer supporting the import from XMI．

LTL and invariant properties have been extended to allow for the use of input signals and next values of state variables．These extensions do not add any expressive power to the language，but facilitates the writing of properties from the user＇s point of view．Internally，each state formula containing a reference to an input or next signal is replaced by a corresponding monitor allowing the reuse of off－the－shelf verification engines．

NUXMV also provides several model transformation techniques aiming to reduce the state space of the design． It uses static analysis techniques to extract possible values for variables，and then re－encode the design using such
information (e.g. using a bit-vector at 32 bit to store 2 values can be re-encoded with just one Boolean variable). These techniques are complemented with others aiming at simplifying the model through constants and free inputs propagation $\left[\mathrm{AFF}^{+} 07\right]$.

Finally, in NUXMV we remove the NUSMV limitation to have bit vectors with less than 64 bits only.

### 1.4 Differences with NUSMV

In this section we summarize the main differences at user level with NUSMV.
As far as the input language of NUXMV is concerned, we introduce two new types for state and input variables. Namely, real and integer (See sections 2.1.5, 2.1.4, and 2.3.1 for details). This enables the user to model the specification of infinite-state transition systems. We add new constructs for specifying the predicates to use in the predicate abstraction based techniques (See sections 2.3.17 and 5.5 for details). nUXMV does not support anymore the keyword process. Indeed, the NUXMV is targeting only finite- and infinite-state synchronous fair transition systems.

We provide the user with all the interactive commands provided by NUSMV (See chapter 4), and we complement them with a set of new commands to use the new features provided by NUXmV (See chapter 5 for a detailed list of the new functionalities provided). In particular, the new commands allows to use the new model checking algorithms for finite-state transition systems, and the new algorithms based on SMT for infinite-state transition systems.

## Structure of this document

This document is structured as follows. First in chapter 2 we describe the syntax and the semantics of the input language of the nUXMV. In chapter 3 we describe how to execute the NUXMV in interactive mode. In chapter 4 we describe the interactive commands inherited from NUSMV, while in chapter 5 we describe the new commands of NUXMV. In chapter 6 we describe the command line switches to execute NUXMV in batch mode (only for finite-state domains).

## Remark

This document is in continuous evolution to better document the features provided by nUXMV.

## Chapter 2

## Input Language of NUXMV

In this chapter we present the syntax and semantics of the input language of NUXMV.
Before going into the details of the language, let us give a few general notes about the syntax. In the syntax notations used below, syntactic categories (non-terminals) are indicated by monospace font, and tokens and character set members (terminals) by bold font. Grammar productions enclosed in square brackets (' []') are optional while a vertical bar ('।') is used to separate alternatives in the syntax rules. Sometimes one of is used at the beginning of a rule as a shorthand for choosing among several alternatives. If the characters I, [ and ] are in bold font, they lose their special meaning and become regular tokens.

In the following, an identifier may be any sequence of characters starting with a character in the set $\{\mathbf{A - Z a - z}\}$ and followed by a possibly empty sequence of characters belonging to the set $\{\mathbf{A - Z a - z 0 - 9 - \$ \# - \}}$. All characters and case in an identifier are significant. Whitespace characters are space ( $\langle$ SPACE $\rangle$ ), tab ( $\langle$ TAB $\rangle$ ) and newline (<RET>). Any string starting with two dashes (' ${ }^{-}-$') and ending with a newline is a comment and ignored by the parser.

The syntax rule for an identifier is:

```
identifier ::
    identifier_first_character
    | identifier identifier_consecutive_character
identifier_first_character :: one of
    A BCD E F G H I J KLM NOP Q R S T U V W X Y Z
    a b c deffgh i j k l m n o p q r s t u v w x y z_
identifier_consecutive_character ::
    identifier_first_character
    | digit
    | one of $ # -
digit :: one of 0 1 2 3 4 5 6 7 8 9
```

An identifier is always distinct from the NUXMV language reserved keywords which are:

A, ABF, ABG, AF, AG, array, ASSIGN, AX, bool, boolean, BU, case, COMPASSION, COMPID, COMPUTE, COMPWFF, CONSTANTS, CONSTRAINT, count, CTLSPEC, CTLWFF, DEFINE, E, EBF, EBG, EF, EG, esac, EX, extend, F, FAIRNESS, FALSE, floor, FROZENVAR, FUN, G, H, IN, in, INIT, init, Integer, integer, INVAR, INVARSPEC, ISA, ITYPE, IVAR, JUSTICE, LTLSPEC, LTLWFF, MAX, MDEFINE, MIN, MIRROR, mod, MODULE, NAME, next, NEXTWFF, O, of, PRED, PREDICATES, PSLSPEC, READ, Real, real, resize, S, self, signed, SIMPWFF, sizeof, SPEC, swconst, T, toint, TRANS, TRUE, U, union, unsigned, uwconst, V, VAR, Word, word, word1, WRITE, X, xnor, xor, $Y, Z$

Note: NUXMV does no longer support the keyword process as only synchronous systems are supported.

To represent various values we will use integer numbers which are any non-empty sequence of decimal digits preceded by an optional unary minus

```
integer_number ::
            - digit
            digit
    | integer_number digit
and symbolic constants which are identifiers
    symbolic_constant :: identifier
```

Examples of integer numbers and symbolic constants are 3, -14, 007, OK, FAIL, waiting, stop. The values of symbolic constants and integer numbers do not intersect.

### 2.1 Types Overview

This section provides an overview of the types that are recognised by nUXmv.

### 2.1.1 Boolean

The boolean type comprises symbolic values FALSE and TRUE.

### 2.1.2 Enumeration Types

An enumeration type is a type specified by full enumerations of all the values that the type comprises. For example, the enumeration of values may be \{stopped, running, waiting, finished\}, $\{2,4,-2,0\}$, $\{$ FAIL , 1, 3, 7, OK\}, etc. All elements of an enumeration have to be unique although the order of elements is not important.

However, in the nUXmV type system, expressions cannot be of actual enumeration types, but of their simplified and generalised versions only. Such generalised enumeration types do not contain information about the exact values constituting the types, but only the flag whether all values are integer numbers, symbolic constants or both. Below only generalised versions of enumeration types are explained.

The symbolic enum type covers enumerations containing only symbolic constants. For example, the enumerations \{stopped, running, waiting\} and \{FAIL, OK\} belong to the symbolic enum type.

There is also a integers-and-symbolic enum type. This type comprises enumerations which contain both integer numbers and symbolic constants, for example, $\{-1,1$, waiting $\},\{0,1,0 K\},\{r u n n i n g$, stopped, waiting, 0$\}$.

Another enumeration type is integer enum. Example of enumerations of integers are $\{2,4,-2,0\}$ and $\{-1,1\}$. In the NUXMV type system an expression of the type integer enum is always converted to the type integer. For explaining the type of expression we will always use the type integer instead of integer enum.

Enumerations cannot contain any boolean value (i.e. $\{$ FALSE, TRUE\}). boolean type must be declared as boolean.

To summarise, we actually deal only with two enumeration types: symbolic enum and integers-andsymbolic enum. These types are distinguishable and have different operations allowed on them.

### 2.1.3 Word

The unsigned word[•] and signed word[•] types are used to model vector of bits (booleans) which allow bitwise logical and arithmetic operations (unsigned and signed, respectively). These types are distinguishable by their width. For example, type unsigned word[3] represents vector of three bits, which allows unsigned operations, and type signed word[7] represents vector of seven bits, which allows signed operations.

When values of unsigned word[ N ] are interpreted as integer numbers the bit representation used is the most popular one, i.e. each bit represents a successive power of 2 between 0 (bit number 0 ) and $2^{N-1}$ (bit number $N-1$ ). Thus unsigned word $[\mathrm{N}]$ is able to represent values from 0 to $2^{N}-1$.

The bit representation of signed word $[\mathrm{N}]$ type is "two's complement", i.e. it is the same as for unsigned word $[\mathrm{N}]$ except that the highest bit (number $N-1$ ) has value $-2^{N-1}$. Thus the possible value for signed word $[\mathrm{N}]$ are from $-2^{N-1}$ to $2^{N-1}-1$.

### 2.1.4 Integer

The domain of the integer type is any Whole Number, positive or negative.
Although the integer type is used to represent integer enum type when explaining the nUXMV type system, there are important differences which are needed to keep in mind. First, using integer is not allowed in certain Model Checking engines and algorithms. Second, at the moment, there are implementation-dependent constraints on the integer enum type, as integer numbers can only be in the range $-2^{32}+1$ to $2^{32}-1$ (more accurately, these values are equivalent to the $\mathrm{C} / \mathrm{C}++$ macros INT_MIN +1 and INT_MAX).

### 2.1.5 Real

The domain of the real type is the Rational Numbers.

### 2.1.6 Array

Arrays are declared with a lower and upper bound for the index, and the type of the elements in the array. For example,
array 0.. 3 of boolean
array 10.. 20 of $\{O K, y, z\}$
array $1 . .8$ of array $-1 . .2$ of unsigned word[5]

The type array 1.. 8 of array $-1 . .2$ of unsigned word[5] means an array of 8 elements (from 1 to 8 ), each of which is an array of 4 elements (from -1 to 2 ) that are 5 -bit-long unsigned words.

Array subtype is the immediate subtype of an array type. For example, subtype of array $1 . .8$ of array $-1 . .2$ of unsigned word[5] is array $-1 . .2$ of unsigned word[5] which has its own subtype unsigned word[5].
array types are incompatible with set type, i.e. array elements cannot be of set type.
Expression of array type can be constructed with array DEFINE (see 2.3.3) or variables of array type (see 2.3.1).

### 2.1.7 Set Types

set types are used to identify expressions representing a set of values. There are four set types: boolean set, integer set, symbolic set, integers-and-symbolic set. The set types can be used in a very limited number of ways. In particular, a variable cannot be of a set type. Only range constant and union operator can be used to create an expression of a set type, and only in, case, ( $\bullet \bullet: \bullet$ ), and assignment ${ }^{1}$ expressions can have imediate operands of a set type.

Every set type has a counterpart among other types. In particular,
the counterpart of a boolean set type is boolean,
the counterpart of a integer set type is integer,
the counterpart of a symbolic set type is symbolic enum,
the counterpart of a integers-and-symbolic set type is integers-and-symbolic enum.
Some types such as unsigned word[॰], signed word[॰], and real do not have a set type counterpart.

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### 2.1.8 Type Order

Figure 2.1 depicts the order existing between types in nUXMV.


Figure 2.1: The ordering on the types in nuXmv
It means, for example, that integer is less than integers-and-symbolic enum and less than real; symbolic enum is less than integers-and-symbolic enum, etc. The unsigned word[•] and signed word $[\bullet]$ types are not comparable with any other type or between each other. Any type is equal to itself.

Note that enumerations containing only integer numbers have the type integer.
For 2 arrays types array N1..M1 of subtype1 and array N2 . M2 of subtype 2 the first type is less then the second one if and only if $\mathrm{N} 1=\mathrm{N} 2, \mathrm{M} 1=\mathrm{M} 2$ and type subtype1 is less than subtype2.

### 2.2 Expressions

In NUXMV all expressions are typed and there are constraints on the type of operands. An expression that violates the type system will be considered erroneous, and will raise a type error.

To maintain backward compatibility with old versions of NUSMV, there is a system variable called backward_compatibility (and a correponding -old command line option) that disables a few new features of NUSMV to keep backward compatibility with old version of NUSMV. In particular, if this system variable is set then type violations caused by expressions of old types (i.e. enumeration type, boolean and integer) will be ignored by the type checker, instead, warnings will be printed out. See the NUSMV user manual [CCCJ ${ }^{+}$10] for further information.

If additionally, the system variable type_checking_warning_on is unset, then even these warnings will not be printed out.

### 2.2.1 Implicit Type Conversion

In some expressions operands may be converted from one type to its set type counterpart (see 2.1.7). For example, integer can be converted to integer set type.

Note: In old version of NUSMV, implicit type conversion from integer to boolean (and vice-versa) was performed. Since NUSMV version 2.5.1, and thus also in NUXMV, implicit integer <-> boolean type conversion is no longer supported, and explicit cast operators have to be used.
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### 2.2.2 Constant Expressions

A constant can be a boolean, integer, symbolic, word or range constant.

```
constant ::
    boolean_constant
    | symbolic_constant
    | integer_constant
    | real_constant
    word_constant
    | range_constant
```


## Boolean Constant

A boolean constant is one of the symbolic values FALSE and TRUE. The type of a boolean constant is boolean.

```
boolean_constant :: one of
    FALSE TRUE
```


## Symbolic Constant

A symbolic constant is syntactically an identifier and indicates a unique value.
symbolic_constant : : identifier
The type of a symbolic constant is symbolic enum. See Section 2.3.14 [Namespaces], page 33 for more information about how symbolic constants are distinguished from other identifiers, i.e. variables, defines, etc.

## Integer Constant

An integer constant is an integer number. The type of an integer constant is integer.
integer_constant :: integer_number

## Real Constant

A real constant is an real number. The type of a real constant is real.
real_constant :: real_number
Definition of real number allows for different representations, namely floating point, fractional and exponential. Some examples:

| float | 123.456 |
| :--- | :--- |
| fractional | $\mathrm{F}^{\prime} 123 / 456$ |
| fractional | $\mathrm{f}^{\prime} 123 / 456$ |
| exponential | 123 e 4 |
| exponential | 123.456 e 7 |
| exponential | 123.456 E 7 |
| exponential | $123.456 \mathrm{E}-7$ |

## Word Constant

Word constant begins with digit 0 , followed by optional character $u$ (unsigned) or $s$ (signed) and one of the characters b/B (binary), o/o (octal), d/D (decimal) or h/H (hexadecimal) which gives the base that the actual constant is in. Next comes an optional decimal integer giving the number of bits, then the character _, and lastly the constant value itself. Assuming N is the width of the constant the type of a word constant is signed word[ N ] if character $s$ is provided, and unsigned word[ N ] otherwise. For example:

```
0sb5_10111 has type signed word[5]
0uo6_37 has type unsigned word[6]
0d11_9 has type unsigned word[11]
0sh12_a9 has type signed word[12]
```

The number of bits can be skipped, in which case the width is automatically calculated from the number of digits in the constant and its base. It may be necessary to explicitly give leading zeroes to make the type correct - the following are all equivalent declarations of the integer constant 11 as a word of type unsigned word[8]:

```
Oud8_11
Oub8_1011
0b_00001011
0h_0b
Oh8_b
```

The syntactic rule of the word constant is the following:

```
word_constant ::
    O [word_sign_specifier] word_base [word_width] _ word_value
word_sign_specifier :: one of
    u s
word_width ::
    integer_number -- a number greater than zero
word_base ::
    b | B | O | O | d | D | h | H
word_value ::
    hex_digit
    word_value hex_digit
    word_value -
hex_digit :: one of
    O 1 2 3 4 5 6 7 8 9 a b c def A B C D E F
```

Note that

- The width of a word must be a number strictly greater than 0 .
- Decimal word constants must be declared with the width specifier, since the number of bits needed for an expression like 0d_019 is unclear.
- Digits are restricted depending on the base the constant is given in.
- Digits can be separated by the underscore character (" -") to aid clarity, for example 0b_0101_1111_1100 which is equivalent to 0b_010111111100.
- For a given width $N$ the value of a constant has to be in range $0 \ldots 2^{N}-1$. For decimal signed words (both $s$ and $d$ are provided) the value of a constant has to be in range $0 \ldots 2^{N-1}$.
- The number of bits in word constant has no longer the implementation limit of being 64 bits at most. In nUXMV it is possible to define words of arbirtary size.


## Range Constant

A range constant specifies a set of consecutive integer numbers. For example, a constant $-1 . .5$ indicates the set of numbers $-1,0,1,2,3,4$ and 5 . Other examples of range constant can be 1..10, $-10 \ldots-10$, 1..300. The syntactic rule of the range constant is the following:

```
range_constant ::
    integer_number .. integer_number
```

with an additional constraint that the first integer number must be less than or equal to the second integer number. The type of a range constant is integer set.

### 2.2.3 Basic Expressions

A basic expression is the most common kind of expression used in NUXMV (as it is also the case in NUSMV).

```
basic_expr ::
    constant
    | variable_identifier
    | define_identifier
    ( basic_expr )
    ! basic_expr
    basic_expr & basic_expr -- logical or bitwise AND
    basic_expr | basic_expr -- logical or bitwise OR
    basic_expr xor basic_expr
    basic_expr xnor basic_expr
    basic_expr -> basic_expr
    basic_expr <-> basic_expr
    basic_expr = basic_expr
    basic_expr != basic_expr
    basic_expr < basic_expr
    basic_expr > basic_expr
    basic_expr <= basic_expr
    basic_expr >= basic_expr
    - basic_expr
    basic_expr + basic_expr
    basic_expr - basic_expr
    basic_expr * basic_expr
    basic_expr / basic_expr
    basic_expr mod basic_expr
    basic_expr >> basic_expr
    basic_expr << basic_expr
    basic_expr [ index ] -- index subscript
    basic_expr [ basic_expr : basic_expr ]
                            -- word bits selection
    basic_expr :: basic_expr -- word concatenation
    word1 ( basic_expr ) -- boolean to unsigned word[1] conversion
    bool ( basic_expr ) -- unsigned word[1] and int to boolean conversion
    toint ( basic_expr ) -- word and boolean to integer constant conversion
    count ( basic_expr_list ) -- count of true boolean expressions
    swconst ( basic_expr , basic_expr )
                            -- integer to signed word constant conversion
    uwconst ( basic_expr, basic_expr )
    -- integer to unsigned word constant conversion
    signed ( basic_expr ) -- unsigned word to signed word conversion
    unsigned ( basic_expr ) -- signed word to unsigned word conversion
    sizeof ( basic_expr ) -- word size as an integer
    floor ( basic_expr ) -- from a real to an integer
    extend ( basic_expr , basic_expr)
-- word width extension
```

```
    | resize ( basic_expr , basic_expr)
    -- word width resize
    basic_expr union basic_expr -- union of set expressions
    { set_body_expr } -- set expression
    basic_expr in basic_expr -- inclusion in a set expression
    basic_expr ? basic_expr : basic_expr
    -- if-then-else expression
    case_expr -- case expression
    basic_next_expr -- next expression
basic_expr_list ::
    basic_expr
    | basic_expr_list , basic_expr
```

The order of parsing precedence for operators from high to low is:

```
[ ] , [ : ]
!
::
- (unary minus)
* / mod
+ -
<< >>
union
in
= != < > <= >=
&
| xor xnor
(\bullet ? \bullet : \bullet)
<->
->
```

Operators of equal precedence associate to the left, except $->$ that associates to the right. The constants and their types are explained in Section 2.2.2 [Constant Expressions], page 11.

## Variables and Defines

A variable_identifier and define_identifier are expressions which identify a variable or a define, respectively. Their syntax rules are:

```
define_identifier :: complex_identifier
variable_identifier :: complex_identifier
```

The syntax and semantics of complex_identifiers are explained in Section 2.3.12 [References to Module Components], page 32. All defines and variables referenced in expressions should be declared. All identifiers (variables, defines, symbolic constants, etc) can be used prior to their definition, i.e. there is no constraint on order such as in C where a declaration of a variable should always be placed in text above the variable use. See more information about define and variable declarations in Section 2.3.2 [DEFINE Declarations], page 26 and Section 2.3.1 [Variable Declarations], page 23.

A define is a kind of macro. Every time a define is met in expressions, it is substituted by the expression associated with this define. Therefore, the type of a define is the type of the associated expression in the current context.
variable_identifier represents state, input, and frozen variables. The type of a variable is specified in its declaration. For more information about variables, see Section 2.3 [Definition of the FSM], page 22, Section 2.3.1 [State Variables], page 24, Section 2.3.1 [Input Variables], page 24, and Section 2.3.1 [Frozen Variables], page 25 . Since a symbolic constant is syntactically indistinguishable from variable_identifiers and define_identifiers, a symbol table is used to distinguish them from each other.

## Parentheses

Parentheses may be used to group expressions. The type of the whole expression is the same as the type of the expression in the parentheses.

## Logical and Bitwise !

The signature of the logical and bitwise NOT operator ! is:

```
! : boolean }->\mathrm{ boolean
```

    : unsigned word[ N\(] \rightarrow\) unsigned word \([\mathrm{N}]\)
    : signed word \([\mathrm{N}] \rightarrow\) signed word \([\mathrm{N}]\)
    This means that the operation can be applied to boolean, unsigned word $[\bullet]$ and signed word $[\bullet]$ operands. The type of the whole expression is the same as the type of the operand. If the operand is not boolean, unsigned word $[\bullet]$ or signed word $[\bullet]$ then the expression violates the type system and nUXMV will throw an error.

## Logical and Bitwise \&, I, xor, xnor, ->, <->

Logical and bitwise binary operators \& (AND), । (OR), xor (exclusive OR), xnor (negated exclusive OR), $\rightarrow$ (implies) and <-> (if and only if) are similar to the unary operator !, except that they take two operands. Their signature is:

```
\&, \(\mid\), xor, xnor, \(\rightarrow\), <-> : boolean * boolean \(\rightarrow\) boolean
    : unsigned word \([\mathrm{N}]\) * unsigned word \([\mathrm{N}] \rightarrow\) unsigned word \([\mathrm{N}]\)
    : signed word[ N\(]\) * signed word \([\mathrm{N}] \rightarrow\) signed word \([\mathrm{N}]\)
```

the operands can be of boolean, unsigned word $[\bullet]$ or signed word $[\bullet]$ type, and the type of the whole expression is the type of the operands. Note that both word operands should have the same width.

## Equality (=) and Inequality ( $!=$ )

The operators $=$ (equality) and $!=$ (inequality) have the following signature:

```
\(=,!=\quad\) : boolean \(*\) boolean \(\rightarrow\) boolean
    integer * integer \(\rightarrow\) boolean
    : integer * real \(\rightarrow\) boolean
    : real * integer \(\rightarrow\) boolean
    : symbolic enum * symbolic enum \(\rightarrow\) boolean
    : integers-and-symbolic enum * integers-and-symbolic enum \(\rightarrow\) boolean
    : unsigned word[ N ] * unsigned word \([\mathrm{N}] \rightarrow\) boolean
    : signed word[ N ] * signed word[N] \(\rightarrow\) boolean
```

No implicit type conversion is performed. For example, in the expression
TRUE = 5
the left operand is of type boolean and the right one is of type integer. Though the signature of the operation does not have a boolean * integer rule, the expression is not correct, because no implicit type conversion will be performed. One can use the toint or the bool for explicit casts.
For example:

```
toint (TRUE) = 5
or
TRUE = bool(5)
```

This is also true if one of the operands is of type unsigned word[1] and the other one is of the type boolean. Explicit cast must be used (e.g. using word1 or bool)

## Relational Operators >, <, >=, <=

The relational operators $>$ (greater than),$<$ (less than), $>=($ greater than or equal to) and $<=$ (less than or equal to) have the following signature:

```
>, <, >=, <= : integer * integer \(\rightarrow\) boolean
    : integer * real \(\rightarrow\) boolean
    : real * integer \(\rightarrow\) boolean
    : unsigned word \([\mathrm{N}] *\) unsigned word \([\mathrm{N}] \rightarrow\) boolean
    : signed word[N] * signed word \([\mathrm{N}] \rightarrow\) boolean
```


## Arithmetic Operators +, -, *, /

The arithmetic operators + (addition), (unary negation or binary subtraction), * (multiplication) and / (division) have the following signature:

```
\(+,-, *, /:\) integer \(*\) integer \(\rightarrow\) integer
    : integer * real \(\rightarrow\) real
    : real * integer \(\rightarrow\) real
    : unsigned word \([\mathrm{N}] *\) unsigned word \([\mathrm{N}] \rightarrow\) unsigned word[ N\(]\)
    : signed word[N] * signed word[ N\(] \rightarrow\) signed word[N]
- (unary) : integer \(\rightarrow\) integer
    : real \(\rightarrow\) real
    : unsigned word \([\mathrm{N}] \rightarrow\) unsigned word[N]
    : signed word[N] \(\rightarrow\) signed word[N]
```

Before checking the expression for being correctly typed, the implicit type conversion can be applied to one of the operands. If the operators are applied to unsigned word[ N$]$ or signed word $[\mathrm{N}]$ type, then the operations are performed modulo $2^{N}$.

The result of the / operator is the quotient from the division of the first operand by the second. The result of the / operator is the algebraic quotient with any fractional part discarded (this is often called "truncation towards zero"). If the quotient $a / b$ is representable, the expression $(a / b) * b+(a \bmod b)$ shall equal $a$. If the value of the second operand is zero, the behavior is undefined and an error is thrown by nUXMV. The semantics is equivalent to the corresponding one of $\mathrm{C} / \mathrm{C}++$ languages.

Similarly to NUSMV, we adopt this new semantics for the division / operator. We refer to the NUSMV user manual $\left[\mathrm{CCCJ}^{+} 10\right]$ for more details on this respect.

## Remainder Operator mod

The result of the mod operator is the algebraic remainder of the division. If the value of the second operand is zero, the behavior is undefined and an error is thrown by nUXMV.

The signature of the remainder operator is:

```
mod : integer * integer }->\mathrm{ integer
    : unsigned word[N] * unsigned word[N] }->\mathrm{ unsigned word[N]
    : signed word[N] * signed word[N] }->\mathrm{ signed word[N]
```

The semantics of mod operator is equivalent to the corresponding operator \% of $\mathrm{C} / \mathrm{C}++$ languages. Thus if the quotient $a / b$ is representable, the expression $(a / b) * b+(a \bmod b)$ shall equal $a$.

Note: in older versions of NUSMV $(\leq 2.4 .0)$ the semantics of quotient and remainder were different. Having the division and remainder operators / and mod be of the current, i.e. C/C++'s, semantics the older semantics of division was given by the formula:

IF (a mod b < 0) THEN ( $\mathrm{a} / \mathrm{b}-1$ ) ELSE ( $\mathrm{a} / \mathrm{b}$ )
and the semantics of remainder operator was given by the formula:
IF $(\mathrm{a} \bmod \mathrm{b}<0)$ THEN $(\mathrm{a} \bmod \mathrm{b}+\mathrm{b})$ ELSE $(\mathrm{a} \bmod \mathrm{b})$
Note that in both interpretations the equation $(a / b) * b+(a \bmod b)=a$ holds. For example, in the current

```
version of NUXMV the following holds
    \(7 / 5=1 \quad 7 \bmod 5=2\)
    \(-7 / 5=-1 \quad-7 \bmod 5=-2\)
    \(7 /-5=-1 \quad 7 \bmod -5=2\)
    \(-7 /-5=1 \quad-7 \bmod -5=-2\)
```

whereas in the old semantics the equations were
$7 / 5=1 \quad 7 \bmod 5=2$
$-7 / 5=-2 \quad-7 \bmod 5=3$
$7 /-5=-1 \quad 7 \bmod -5=2$
$-7 /-5=0 \quad-7 \bmod -5=-7$

When supplied, the command line option -old_div_op switches the semantics of division and remainder to the old one.

## Shift Operators <<, >>

The signature of the shift operators is:

```
<<, >> : unsigned word[N] * integer }->\mathrm{ unsigned word[N]
    : signed word[N] * integer }->\mathrm{ signed word[N]
    : unsigned word[N] * unsigned word[M] }->\mathrm{ unsigned word[N]
    : signed word[N] * unsigned word[M] }->\mathrm{ signed word[N]
```

Before checking the expression for being correctly typed, the right operand can be implicitly converted from boolean to integer type.

Left shift << (right shift >>) operation shifts to the left (right) the bits of the left operand by the number specified in the right operand. A shift by N bits is equivalent to N shifts by 1 bit. A bit shifted behind the word bound is lost. During shifting a word is padded with zeros with the exception of the right shift for signed word $[\bullet]$, in which case a word is padded with its highest bit. For instance,

$$
\begin{aligned}
& \text { Oub4_0101 } \ll 2 \text { is equal to } \\
& \text { Oub4_0100 } \ll 1 \text { is equal to } \\
& \text { Oub4_1000 } \ll 0 \text { is equal to } \\
& \text { Oub4_1000 and }
\end{aligned}
$$

0sb3_1011 >> 2 is equal to
0sb3_1110 >> 1 is equal to
0sb3_1111 >> 0 is equal to
0sb3_1111

It has to be remarked that the shifting requires the right operand to be greater or equal to zero and less then or equal to the width of the word it is applied to. NUXMV raises an error if a shift is attempted that does not satisfy this restriction.

## Index Subscript Operator [ ]

The index subscript operator extracts one element of an array in the typical fashion. On the left of [ ] there must be an expression of array type. The index expression in the brackets has to be an expression of integer or word[•] type with value greater or equal to lower bound and less or equal to the upper bound of the array. The signature of the index subscript operator is:

> [ ]: array N..M of subtype $*$ word $[\mathrm{N}] \rightarrow$ subtype
> $\quad:$ array N..M of subtype $*$ integer $\rightarrow$ subtype

For example, for below declarations ${ }^{2}$ :

```
MODULE main
    VAR a : array -1 .. 4 of array 1 .. 2 of boolean;
    DEFINE d := [[12, 4], [-1,2]];
    VAR r : 0..1;
```

expressions $a[-1]$, $a[0][r+1]$ and $d[r][1]$ are valid whereas $a[0]$, $a[0][r]$ and $d[0][r-1]$ will raise an out of bound error.

[^1]
## Bit Selection Operator [ : ]

The bit selection operator extracts consecutive bits from a unsigned word[•] or signed word[•] expression, resulting in a new unsigned word $[\bullet]$ expression. This operation always decreases the width of a word or leaves it intact. The expressions in the brackets have to be integer constants which specify the high and low bound. The high bound must be greater than or equal to the low bound. The bits count from 0 . The result of the operations is unsigned word[•] value consisting of the consecutive bits beginning from the high bound of the operand down to, and including, the low bound bit. For example, 0sb7_1011001[4:1] extracts bits 1 through 4 (including 1st and 4th bits) and is equal to 0ub4_1100. 0ub3_101[0:0] extracts bit number 0 and is equal to 0 ub1_1.

The signature of the bit selection operator is:

```
[ : ] : unsigned word[ N ] \(*\) integer \(_{h} *\) integer \(_{l} \rightarrow\) unsigned word[integer \({ }_{h}\) - integer \(_{l}+1\) ]
        : signed word \([\mathrm{N}] *\) integer \(_{h} *\) integer \(_{l} \rightarrow\) unsigned word[integer \(_{h}-\) integer \(\left._{l}+1\right]\)
where \(0 \leq\) integer \(_{l} \leq\) integer \(_{h}<\mathrm{N}\)
```


## Word Concatenation Operator : :

The concatenation operator joins two words (unsigned word[•] or signed word[•] or both) together to create a larger unsigned word[•] type. The operator itself is two colons (: :), and its signature is as follows:

$$
::: \operatorname{word}[\mathrm{M}] * \text { word }[\mathrm{N}] \rightarrow \text { unsigned word }[\mathrm{M}+\mathrm{N}]
$$

where word $[\mathrm{N}]$ is unsigned word $[\mathrm{N}]$ or signed word $[\mathrm{N}]$. The left-hand operand will make up the upper bits of the new word, and the right-hand operand will make up the lower bits. The result is always unsigned word[•]. For example, given the two words w1 $:=0 u b 4 \_1101$ and $w 2:=0 s b 2 \_00$, the result of w1::w2 is 0ub6_110100.

## Extend Word Conversions

extend operator increases the width of a word by attaching additional bits on the left. If the provided word is unsigned then zeros are added, otherwise if the word is signed the highest (sing) bit is repeated corresponding number of times.

The signature of the operator is:
extend: unsigned word[ N ] $*$ integer $\rightarrow$ unsigned word[ $\mathrm{N}+$ +integer ]

$$
: \text { signed word }[\mathrm{N}] * \text { integer } \rightarrow \text { signed word }[\mathrm{N}+\text { integer }]
$$

For example:

$$
\begin{aligned}
& \text { extend(0ub3_101, 2) }=0 \text { ubb5_00101 } \\
& \text { extend(0sb3_101, })=0 \text { sb5_11101 } \\
& \text { extend(0sb3_011, 2) }=0 \text { sb5_00011 }
\end{aligned}
$$

Note that the right operand of extend has to be an integer constant greater or equal to zero.

## Resize Word Conversions

resize operator provides a more comfortable way of changing the word of a width. The behavior of this operator can be described as follows:

Let w be a M bits unsigned word $[\bullet]$ and N be the required width: if $\mathrm{M}=\mathrm{N}$, w is returned unmodified; if N is less than $M$, bits in the range $[N-1: 0]$ are extracted from $w$; if $N$ is greater than $M$, $w$ is extended of $(N-M)$ bits up to required width, padding with zeroes.

Let w be a M bits signed word $[\bullet]$ and N be the required width: if $\mathrm{M}=\mathrm{N}$, w is returned unmodified; if N is less than M , bits in the range $[\mathrm{N}-2: 0]$ are extracted from w , while $\mathrm{N}-1$-ith bit is forced to preserve the value of the original sign bit of w ( $\mathrm{M}-1$-ith bit); if N is greater than M , w is extended of $(\mathrm{N}-\mathrm{M})$ bits up to required width, extending sign bit.

The signature of the operator is:

```
resize : unsigned word[•] * integer \(\rightarrow\) unsigned word[integer ]
    : signed word[•] * integer \(\rightarrow\) signed word[integer ]
```


## Set Expressions

The set expression is an expression defining a set of boolean, integer and symbolic enum values. A set expression can be created with the union operator. For example, 1 union 0 specifies the set of values 1 and 0 . One or both of the operands of union can be sets. In this case, union returns a union of these sets. For example, expression ( 1 union 0 ) union -3 specifies the set of values 1,0 and -3 .

Note that there cannot be a set of sets in NUXMV . Sets can contain only singleton values, but not other sets.
The signature of the union operator is:

```
union : boolean set * boolean set }->\mathrm{ boolean set
    : integer set * integer set }->\mathrm{ integer set
    : symbolic set * symbolic set }->\mathrm{ symbolic set
    : integers-and-symbolic set * integers-and-symbolic set
    integers-and-symbolic set
```

Before checking the expression for being correctly typed, if it is possible, both operands are converted to their counterpart set types ${ }^{3}$, which virtually means converting individual values to singleton sets. Then both operands are implicitly converted to a minimal type that covers both operands. If after these manipulations the operands do not satisfy the signature of union operator, an error is raised by NUXMV.

There is also another way to write a set expression by enumerating all its values between curly brackets. The syntactic rule for the values in curly brackets is:

```
set_body_expr ::
    basic_expr
    | set_body_expr , basic_expr
```

Enumerating values in curly brackets is semantically equivalent to writing them connected by union operators. For example, expression $\{\exp 1, \exp 2, \exp 3\}$ is equivalent to $\exp 1$ union $\exp 2$ union exp3. Note that according to the semantics of union operator, expression $\{\{1,2\},\{3,4\}\}$ is equivalent to $\{1,2,3,4\}$, i.e. there is no actually set of sets.

Set expressions can be used only as operands of union and in operations, as the right operand of case and as the second and the third operand of ( $\bullet \bullet: \bullet)$ expressions and assignments. In all other places the use of set expressions is prohibited.

## Inclusion Operator in

The inclusion operator 'in' tests the left operand for being a subset of the right operand. If either operand is a number or a symbolic value instead of a set, it is coerced to a singleton set.

The signature of the in operator is:

```
in : boolean set * boolean set }->\mathrm{ boolean
    : integer set * integer set }->\mathrm{ boolean
    : symbolic set * symbolic set }->\mathrm{ boolean
    : integers-and-symbolic set * integers-and-symbolic set }->\mathrm{ boolean
```

Similar to union operation, before checking the expression for being correctly typed, if it is possible, both operands are converted to their counterpart set types ${ }^{4}$. Then, if required, implicit type conversion is carried out on one of the operands.

[^2]
## Case Expressions

A case expression has the following syntax:

```
case_expr :: case case_body esac
case_body ::
    basic_expr : basic_expr ;
    | case_body basic_expr : basic_expr ;
```

A case_expr returns the value of the first expression on the right hand side of ' $:$ ', such that the corresponding condition on the left hand side evaluates to tRUE. For example, the result of the expression

```
case
    left_expression_1 : right_expression_1 ;
    left_expression_2 : right_expression_2 ;
    left_expression_N : right_expression_N ;
esac
```

is right_expression_k such that for all $i$ from 0 to $k-1$, left_expression_i is FALSE, and left_expression_k is TRUE. It is an error if all expressions on the left hand side evaluate to FALSE.

The type of expressions on the left hand side must be boolean. If one of the expression on the right is of a set type then, if it is possible, all remaining expressions on the right are converted to their counterpart set types ${ }^{5}$. The type of the whole expression is such a minimal type ${ }^{6}$ that all of the expressions on the right (after possible convertion to set types) can be implicitly converted to this type. If this is not possible, nUXMV throws an error.

Note: Prior to version 2.5.1, using 1 as left_expression_N was pretty common, e.g:

```
case
    cond1 : expr1;
    cond2 : expr2;
    .
    1 : exprN; -- otherwise
esac
```

Since version 2.5.1 integer values are no longer implicitly casted to boolean, and 1 has to be written as TRUE instead.

## If-Then-Else expressions

In certain cases, the syntax described above may look a bit awkard. In simpler cases, it is possible to use the alternative, terser, ( $\bullet \bullet: \bullet)$ expression. This construct is defined as follows:

```
cond_expr ? basic_expr1 : basic_expr2
```

This expression evaluates to basic_expr1 if the condition in cond_expr evaluates to true, and to basic_expr2 otherwise. Therefore, the expressions cond1 ? expr1 : expr2 and case cond1 : expr1; TRUE : expr2; esac are equivalent.

## Basic Next Expression

Next expressions refer to the values of variables in the next state. For example, if a variable $\mathbf{v}$ is a state variable, then next (v) refers to that variable $\mathbf{v}$ in the next time step. A next applied to a complex expression is a shorthand method of applying next to all the variables in the expressions recursively. Example: next ( $(1+$ a) + b) is equivalent to ( $1+\operatorname{next}(\mathrm{a}))+\operatorname{next}(\mathrm{b})$. Note that the next operator cannot be applied twice, i.e. next (next (a)) is not allowed.

The syntactic rule is:

[^3]$\underset{\substack{\text { FONDAZOONE } \\ \text { BRINO KESSER }}}{ }$

```
basic_next_expr :: next ( basic_expr )
```

A next expression does not change the type.

## Count Operator

The count operator counts the number of expressions which are true. The count operator is a syntactic sugar for

```
toint (bool_expr1) +
toint (bool_expr2) +
... +
toint (bool_exprN)
```

This operator has been introduced in version 2.5.1, to simplify the porting of those models which exploited the implicit casting of integer to boolean to encoding e.g. predicates like:
(b0 + b1 + ... +bN ) < 3 -- at most two bits are enabled Since version 2.5.1, this expression can be written as:

```
count (b0 + b1 + ... + bN) < 3
```


### 2.2.4 Simple and Next Expressions

Simple_expressions are expressions built only from the values of variables in the current state. Therefore, the simple_expression cannot have a next operation inside and the syntax of simple_expressions is as follows:

```
simple_expr :: basic_expr
```

with the alternative basic_next_expr not allowed. Simple_expressions can be used to specify sets of states, for example, the initial set of states. The next_expression relates current and next state variables to express transitions in the FSM. The next_expression can have next operation inside, i.e.

```
next_expr :: basic_expr
```

with the alternative basic_next_expr allowed.

### 2.2.5 Type conversion operators

## Integer conversion operator

toint converts an unsigned word[•] constant or a signed word[॰] constant, or a boolean expression to an integer representing its value. Also integer expressions are allowed, but no action is performed. The signature of this conversion operator is:

```
toint : integer }->\mathrm{ integer
toint: boolean }->\mathrm{ integer
toint: unsigned word[\bullet] }->\mathrm{ integer
toint : signed word[\bullet] }->\mathrm{ integer
```


## floor convertion operator

The operator floor maps a real number to the largest previous integer. If applied to an integer it returns the integer itself. It has the following signature:

```
floor: integer }->\mathrm{ integer
    : real }->\mathrm{ integer
```


## Boolean conversion operator

bool converts unsigned word[1] and any expression of type integer (e.g. $1+2$ ) to boolean. Also boolean expressions are allowed, but no action is perfomed. In case of integer expression, the result of the conversion is FALSE if the expression resolves to 0 , TRUE otherwise. In case of unsigned word[1] expression, the conversion obeys the following table:

$$
\begin{aligned}
& \text { bool }\left(0 \mathrm{ub} 1 \_0\right)=\text { FALSE } \\
& \text { bool }\left(0 \mathrm{ub} 1 \_1\right)=\text { TRUE }
\end{aligned}
$$

## Integer to Word Constants Conversion

swconst, uwconst convert an integer constant into a signed word[•] constant or unsigned word[•] constant of given size respectively. The signature of these conversion operator is:

```
swconst : integer * integer }->\mathrm{ signed word[॰]
uwconst : integer * integer }->\mathrm{ unsigned word[॰]
```


## Word1 Explicit Conversions

word1 converts a boolean to a unsigned word[1]. The signature of this conversion operator is:

```
word1 : boolean }->\mathrm{ unsigned word[1]
```

The conversion obeys the following table:

$$
\begin{aligned}
& \text { word1(FALSE) = 0ub1_0 } \\
& \text { word1(TRUE) = 0ub1_1 }
\end{aligned}
$$

## Unsigned and Signed Explicit Conversions

unsigned converts a signed word[ N$]$ to an unsigned word[ N$]$, while signed performs the opposite operation and converts an unsigned word $[\mathrm{N}]$ to a signed word $[\mathrm{N}]$. Both operations do not change the bit representation of a provided word. The signatures of these conversion operators are:

```
unsigned : signed word[N] }->\mathrm{ unsigned word[N]
signed : unsigned word[N] }->\mathrm{ signed word[N]
```

For example:

$$
\begin{aligned}
& \text { signed(0ub_101) = 0sb_101 } \\
& \text { signed(0ud3_5) = -0sd3_3 } \\
& \text { unsigned(0sb_101) = 0usb_101 } \\
& \text { unsigned(-0sd3_3) = 0ud3_5 }
\end{aligned}
$$

### 2.3 Definition of the FSM

We consider a Finite State Machine (FSM) described in terms of state variables input variables, and frozen variables, which may assume different values in different states; of a transition relation describing how inputs leads from one state to possibly many different states; and of Fairness conditions that describe constraints on the valid paths of the execution of the FSM. In this document, we distinguish among constraints (used to constrain the behavior of a FSM, e.g. a modulo 4 counter increments its value modulo 4), and specifications (used to express properties to verify on the FSM (e.g. the counter reaches value 3).

In the following it is described how these concepts can be declared in the nUXmv language.

### 2.3.1 Variable Declarations

A variable can be an input, a frozen, or a state variable. The declaration of a variable specifies the variable's type with the help of type specifier.

## Type Specifiers

```
A type specifier has the following syntax:
type_specifier ::
        simple_type_specifier
    | module_type_specifier
simple_type_specifier ::
        boolean
        word [ basic_expr ]
        unsigned word [ basic_expr ]
        signed word [ basic_expr ]
        real
        integer
        { enumeration_type_body }
        basic_expr .. basic_expr
        array basic_expr .. basic_expr
            of simple_type_specifier
enumeration_type_body ::
    enumeration_type_value
    | enumeration_type_body , enumeration_type_value
enumeration_type_value ::
    symbolic_constant
    | integer_number
```

There are two kinds of type specifier: a simple type specifier and a module type specifier. The module type specifier is explained later in Section 2.3.11 [MODULE Instantiations], page 31. The simple type specifier comprises boolean type, integer type, enumeration types, unsigned word[•], signed word[ $\bullet \cdot]$ and arrays types.

The boolean type is specified by the keyword boolean. The integer type is specified by the keyword integer, and the The real type is specified by the keyword real.

A enumeration type is specified by full enumeration of all the values the type comprises. For example, possible enumeration type specifiers are $\{0,2,3,-1\},\{1,0, O K\},\{0 K$, FAIL, running $\}$. FALSE and TRUE values cannot be used as enumeration type specifiers The values in the list are enclosed in curly brackets and separated by commas. The values may be integer numbers, symbolic constants, or both. All values in the list should be distinct from each other, although the order of values is not important.

Note, expressions cannot be of the actual enumeration types, but only the simplified versions of enumeration types, such as symbolic enum and integers-and-symbolic enum.

A type specifier can be given by two expressions separated by .. (<TWO DOTS>). The two expressions have both to evaluate to constants integer numbers, and may contain names of defines and module formal parameters. For example, $-1-P 1 \ldots 5+D 1$, where P1 refers to a module formal parameter, and D1 refers to a define. Both P1 and D1 have to be statically evaluable to integer constants.

This is just a shorthand for a enumeration type containing the list of integer numbers from the range given in type specifier. For example, the type specifiers $-1 \ldots 5$ and $\{-1,0,1,2,3,4,5\}$ are equivalent. Note that the evaluated number on the left from the two dots must be less than or equal to the evaluated number on the right.

The unsigned word[•] type is specified by the keywords unsigned word (where unsigned may be skipped) with a basic_expr supplied in square brackets. The expression must be statically evaluable to a constant integer number whose value must be greater than zero. The signed word[•] type is specified in a similar way with the keywords signed word. The purpose of the word types is to offer integer and bitwise arithmetic.

An array type is denoted by a sequence of the keyword array, a basic_expr specifying the lower bound of the array index, two dots . ., a basic_expr specifying the upper bound of the array index, the keyword of, and the type of array's elements. The elements can themselves be arrays. The two bound expressions have to be statically evaluable to constant integer numbers, and may contain names of defines and module formal parameters.

## State Variables

A state of the model is an assignment of values to a set of state and frozen variables. State variables (and also instances of modules) are declared by the notation:

```
var_declaration :: VAR var_list
var_list :: identifier : type_specifier ;
    | var_list identifier : type_specifier ;
```

A variable declaration specifies the identifier of the variables and its type. A variable can take the values only from the domain of its type. In particular, a variable of a enumeration type may take only the values enumerated in the type specifier of the declaration.

## Input Variables

IVAR s (input variables) are used to label transitions of the Finite State Machine. The difference between the syntax for the input and state variables declarations is the keyword indicating the beginning of a declaration:

```
ivar_declaration :: IVAR simple_var_list
simple_var_list ::
    identifier : simple_type_specifier ;
    | simple_var_list identifier : simple_type_specifier ;
```

Another difference between input and state variables is that input variables cannot be instances of modules. The usage of input variables is more limited than the usage of state variables which can occur everywhere both in the model and specifications. Namely, input variables cannot occur in:

- Left-side of assignments. For example all these assignments are not allowed:

```
IVAR i : boolean;
ASSIGN
init(i) := TRUE;
next(i) := FALSE;
```

- InIt statements. For example:

```
IVAR i : boolean;
VAR s : boolean;
INIT i = s
```

- Scope of next expressions. For example:

```
IVAR i : boolean;
VAR s : boolean;
TRANS i -> s-this is allowed
TRANS next(i -> s) - this is NOT allowed
```

- Some specification kinds: CTLSPEC, SPEC, COMPUTE, PSLSPEC. For example:

```
IVAR i : boolean;
VAR s : boolean;
SPEC AF (i -> s) - this is NOT allowed
LTLSPEC F (X i -> s) - this is allowed
INVARSPEC (i -> s) - this is allowed
```


## Frozen Variables

Frozen variables are variables that retain their initial value throughout the evolution of the state machine; this initial value can be constrained in the same ways as for normal state variables. Similar to input variables the difference between the syntax for the frozen and state variables declarations is the keyword indicating the beginning of a declaration:
frozenvar_declaration : : FROZENVAR simple_var_list
The semantics of a frozen variable $f v$ is that of a state variable accompanied by an assignment that keeps its value constant (it is handled more efficiently, though):

```
ASSIGN next(fv) := fv;
```

As a consequence, frozen variables may not have their current and next value set in an ASSIGN statement, i.e. statements such as ASSIGN next (fv) :=expr; and ASSIGN fv := expr; are illegal. Apart from that frozen variables may occur in the definition of the FSM in any place in which a state variable may occur. Some examples are as follows:

- Left-side current and next state assignments are illegal, while init state assignments are allowed:

```
FROZENVAR a : boolean;
FROZENVAR b : boolean;
FROZENVAR c : boolean;
VAR d : boolean;
FROZENVAR e : boolean;
ASSIGN
init(a) := d; -- legal
next(b) := d; -- illegal
c := d; -- illegal
e := a; -- also illegal
```

- INIT, TRANS, INVAR, FAIRNESS, JUSTICE, and COMPASSION statements are all legal. So is the scope of a next expression. For example:

```
-- the following has an empty state space
FROZENVAR a : boolean;
INIT a
INVAR !a
-- alternatively, this has two initial states, deadlocking
FROZENVAR b : boolean;
TRANS next(b) <-> !b
-- and that's just unfair
FROZENVAR c : boolean;
FAIRNESS C
FAIRNESS !c
```

- All kinds of specifications involving frozen variables are allowed, e.g.:

```
FROZENVAR c : boolean;
```

-- True by definition.
SPEC AG ((c -> AG c) \& ((!c) -> AG !c))
-- Here, neither is true.
INVARSPEC C
INVARSPEC ! c
-- False (as above).
LTLSPEC (G F c) \& (G F ! c)

## Examples

Below are examples of state, frozen, and input variable declarations:

```
VAR a : boolean;
FROZENVAR b : 0..1;
IVAR c : {TRUE, FALSE};
```

The variable $a$ is a state variable, $b$ is a frozen variable, and $c$ is an input variable; In the following examples:

```
VAR d : {stopped, running, waiting, finished};
VAR e : {2, 4, -2, 0};
VAR f : {1, a, 3, d, q, 4};
```

the variables $d$, e and $f$ are of enumeration types, and all their possible values are specified in the type specifiers of their declarations.

```
VAR g : unsigned word[3];
VAR h : word[3];
VAR i : signed word[4];
```

The variables $g$ and $h$ are typeunsigned word 3-bits-wide (i.e. unsigned word[3]), and $i$ is an signed word 4-bits-wide (i.e. signed word[4]).

```
VAR j : array -1..1 of boolean;
```

The variable $j$ is an array of boolean elements with indexes $-1,0$ and 1 .

### 2.3.2 define Declarations

In order to make descriptions more concise, a symbol can be associated with a common expression, and a DEFINE declaration introduces such a symbol. The syntax for this kind of declaration is:

```
define_declaration :: DEFINE define_body
define_body :: identifier := next_expr ;
    | define_body identifier := next_expr ;
```

DEFINE associates an identifier on the left hand side of the ':=' with an expression on the right side. A define statement can be considered as a macro. Whenever a define identifier occurs in an expression, the identifier is syntactically replaced by the expression it is associated with. The associated expression is always evaluated in the context of the statement where the identifier is declared (see Section 2.3.15 [Context], page 34 for an explanation of contexts). Forward references to defined symbols are allowed but circular definitions are not, and result in an error. The difference between defined symbols and variables is that while variables are statically typed, definitions are not.

### 2.3.3 Array Define Declarations

It is possible to specify an array expressions. This feature is experimental and currently available only through define declaration. The syntax for this kind of declaration is:

```
array_define_declaration ::
    DEFINE identifier := array_expression ;
array_expression :: [ array_contents ]
    | [ array_expression_list ]
```

```
array_expression_list :: array_expression
    | array_expression , array_expression_list
array_contents :: next_expr , array_contents
    | next_expr
```

Array DEFINE associates an identifier on the left hand side of the ':=' with an array expression. As a normal DEFINE statement an array define is considered as a macro. Whenever an array identifier occurs in an expression, the identifier is syntactically replaced by the array expression it is associated with. As with normal define an array define expression is always evaluated in the context of the statement where the identifier is declared and forward references to defined symbols are allowed but circular definitions are not.

The type of an array expression [exp1, exp2, ..., expN] is array $0 . . \mathrm{N}-1$ of type where type is the least type such that all exp1, exp2, ...expN can be converted to it.

It is not possible to declare asymmetrical arrays. This means that it is forbidden to declare an array with a different number of elements in a dimension. For example, the following code will result in an error:

```
DEFINE
    x := [[1,2,3], [1,2]];
```


### 2.3.4 constants Declarations

CONSTANTS declarations allow the user to explicitly declare symbolic constants that might occur or not within the FSM that is being defined. CONSTANTS declarations are expecially useful in those conditions that require symbolic constants to occur only in DEFINEs body (e.g. in generated models). For an example of usage see also the command write_boolean_model. A constant is allowed to be declared multiple times, as after the first declaration any further declaration will be ignored. CONSTANTS declarations are an extension of the original SMV grammar. They have been integrated in NuSMV and inherited in nUXmv. The syntax for this kind of declaration is:

```
constants_declaration :: CONSTANTS constants_body ;
constants_body :: identifier
    | constants_body , identifier
```


### 2.3.5 init Constraint

The set of initial states of the model is determined by a boolean expression under the INIT keyword. The syntax of an INIT constraint is:
init_constrain : : INIT simple_expr [;]
Since the expression in the INIT constraint is a simple_expression, it cannot contain the next () operator. The expression also has to be of type boolean. If there is more than one INIT constraint, the initial set is the conjunction of all of the INIT constraints.

### 2.3.6 invar Constraint

The set of invariant states can be specified using a boolean expression under the INVAR keyword. The syntax of an INVAR constraint is:

```
invar_constraint :: INVAR simple_expr [;]
```

Since the expression in the INVAR constraint is a simple_expression, it cannot contain the next () operator. If there is more than one INVAR constraint, the invariant set is the conjunction of all of the INVAR constraints.

### 2.3.7 trans Constraint

The transition relation of the model is a set of current state/next state pairs. Whether or not a given pair is in this set is determined by a boolean expression, introduced by the trans keyword. The syntax of a TRANS constraint is:

```
trans_constraint :: TRANS next_expr [;]
```

It is an error for the expression to be not of the boolean type. If there is more than one TRANS constraint, the transition relation is the conjunction of all of TRANS constraints.

### 2.3.8 assign Constraint

An assignment has the form:

```
assign_constraint :: ASSIGN assign_list
assign_list :: assign ;
    | assign_list assign ;
assign ::
    complex_identifier := simple_expr
    | init ( complex_identifier ) := simple_expr
    | next ( complex_identifier ) := next_expr
```

On the left hand side of the assignment, identifier denotes the current value of a variable, 'init(identifier)' denotes its initial value, and 'next (identifier)' denotes its value in the next state. If the expression on the right hand side evaluates to a not-set expression such as integer number or symbolic constant, the assignment simply means that the left hand side is equal to the right hand side. On the other hand, if the expression evaluates to a set, then the assignment means that the left hand side is contained in that set. It is an error if the value of the expression is not contained in the range of the variable on the left hand side.

Semantically assignments can be expressed using other kinds of constraints:

```
ASSIGN a := exp; is equivalent to INVAR a in exp;
ASSIGN init(a) := exp; is equivalent to INIT a in exp;
ASSIGN next(a) := exp; is equivalent to TRANS next(a) in exp;
```

Notice that, an additional constraint is forced when assignments are used with respect to their corresponding constraints counterpart: when a variable is assigned a value that it is not an element of its declared type, an error is raised.

The allowed types of the assignment operator are:

```
:= : integer * integer
    : real * integer
    : real * real
    : integer * integer set
    symbolic enum * symbolic enum
    : symbolic enum * symbolic set
    : integers-and-symbolic enum * integers-and-symbolic enum
    : integers-and-symbolic enum * integers-and-symbolic set
    : unsigned word[N] * unsigned word[N]
    : signed word[N] * signed word[N]
```

Before checking the assignment for being correctly typed, the implicit type conversion can be applied to the right operand.

## Rules for assignments

Assignments describe a system of equations that say how the FSM evolves through time. With an arbitrary set of equations there is no guarantee that a solution exists or that it is unique. We tackle this problem by placing certain restrictive syntactic rules on the structure of assignments, thus guaranteeing that the program is implementable.

The restriction rules for assignments are:

- The single assignment rule - each variable may be assigned only once.
- The circular dependency rule - a set of equations must not have "cycles" in its dependency graph not broken by delays (i.e. by a next, see examples below).

The single assignment rule disregards conflicting definitions, and can be formulated as: one may either assign a value to a variable " $x$ ", or to "next ( $x$ )" and "init ( $x$ )", but not both. For instance, the following are legal assignments:

| Example 1 | $\mathrm{x}:=\operatorname{expr} r_{1} ;$ |
| :--- | :--- |
| Example 2 | $\operatorname{init}(\mathrm{x}):=\operatorname{expr} r_{1} ;$ |
| Example 3 | $\operatorname{next}(\mathrm{x}):=\operatorname{expr}_{1} ;$ |
| Example 4 | init $(\mathrm{x}):=\operatorname{expr}_{1} ;$ <br> $\operatorname{next}(\mathrm{x}):=\operatorname{expr}_{2} ;$ |

while the following are illegal assignments:

| Example 1 | $\begin{aligned} & \mathrm{x}:=\operatorname{expr}_{1} ; \\ & \mathrm{x}:=\operatorname{expr}_{2} ; \end{aligned}$ |
| :---: | :---: |
| Example 2 | $\begin{aligned} & \text { init }(x):=\operatorname{expr}_{1} ; \\ & \text { init }(x):=\operatorname{expr}_{2} ; \end{aligned}$ |
| Example 3 | $\begin{aligned} & \mathrm{x}:=\exp r_{1} ; \\ & \text { init }(x):=\operatorname{expr}_{2} ; \end{aligned}$ |
| Example 4 | $\begin{aligned} & \mathrm{x}:=\operatorname{expr}_{1} ; \\ & \text { next }(\mathrm{x}):=\operatorname{expr}_{2} ; \end{aligned}$ |

If we have an assignment like $\mathrm{x}:=\mathrm{y}$; , then we say that x depends on y . A combinatorial loop is a cycle of dependencies not broken by delays. For instance, the assignments:

```
x := y;
y := x;
```

form a combinatorial loop. Indeed, there is no fixed order in which we can compute x and y , since at each time instant the value of $x$ depends on the value of $y$ and vice-versa. We can introduce a "unit delay dependency" using the next () operator.

```
    x := y;
next(y) := x;
```

In this case, there is a unit delay dependency between $x$ and $y$. A combinatorial loop is a cycle of dependencies whose total delay is zero. In nUXMV (as well as in NUSMV) combinatorial loops are illegal. This guarantees that for any set of equations describing the behavior of variable, there is at least one solution. There might be multiple solutions in the case of unassigned variables or in the case of non-deterministic assignments such as in the following example,

```
next(x) := case x = 1 : 1;
    TRUE : {0,1};
    esac;
```


### 2.3.9 fairness Constraints

A fairness constraint restricts the attention only to fair execution paths. When evaluating specifications, the model checker considers path quantifiers to apply only to fair paths.
nUXMV (as well as NUSMV) supports two types of fairness constraints, namely justice constraints and compassion constraints. A justice constraint consists of a formula $f$, which is assumed to be true infinitely often in all the fair paths. In NUXMV, justice constraints are identified by keywords JUSTICE and, for backward compatibility, FAIRNESS. A compassion constraint consists of a pair of formulas ( $\mathrm{p}, \mathrm{q}$ ) ; if property p is true infinitely often in a fair path, then also formula $q$ has to be true infinitely often in the fair path. In nUXMV, compassion constraints are identified by keyword COMPASSION. ${ }^{7}$ Note: If compassion constraints are used, then the model must not contain any input variable. Currently, nUXMV does not enforce this so it is the responsibility of the user to make sure that this is the case.

Fairness constraints are declared using the following syntax (all expressions are expected to be boolean):

```
fairness_constraint ::
    FAIRNESS simple_expr [;]
    JUSTICE simple_expr [;]
    COMPASSION ( simple_expr , simple_expr ) [;]
```

A path is considered fair if and only if it satisfies all the constraints declared in this manner.

### 2.3.10 module Declarations

A module declaration is an encapsulated collection of declarations, constraints and specifications. A module declaration also opens a new identifier scope. Once defined, a module can be reused as many times as necessary. Modules are used in such a way that each instance of a module refers to different data structures. A module can contain instances of other modules, allowing a structural hierarchy to be built. The syntax of a module declaration is as follows:

```
module :: MODULE identifier [( module_parameters )] [module_body]
module_parameters ::
    identifier
    | module_parameters , identifier
module_body ::
            module_element
            | module_body module_element
module_element ::
            var_declaration
    | ivar_declaration
    | frozenvar_declaration
    | define_declaration
    | constants_declaration
    | assign_constraint
    | trans_constraint
    | init_constraint
    | invar_constraint
    | fairness_constraint
    | ctl_specification
    | invar_specification
    | ltl_specification
    | compute_specification
    | isa_declaration
    | pred_declaration
```

[^4]```
| mirror_declaration
```

The identifier immediately following the keyword mODULE is the name associated with the module. Module names have a separate name space in the program, and hence may clash with names of variables and definitions. The optional list of identifiers in parentheses are the formal parameters of the module.

### 2.3.11 mODULE Instantiations

An instance of a module is created using the VAR declaration (see Section 2.3.1 [State Variables], page 24) with a module type specifier (see Section 2.3.1 [Type Specifiers], page 23). The syntax of a module type specifier is:

```
module_type_specifier ::
    identifier [ ( [ parameter_list ] ) ]
parameter_list ::
    next_expr
    | parameter_list , next_expr
```

A variable declaration with a module type specifier introduces a name for the module instance. The module type specifier provides the name of the instantiating module and also a list of actual parameters, which are assigned to the formal parameters of the module. An actual parameter can be any legal next expression (see Section 2.2.4 [Simple and Next Expressions], page 21). It is an error if the number of actual parameters is different from the number of formal parameters. Whenever formal parameters occur in expressions within the module, they are replaced by the actual parameters. The semantic of module instantiation is similar to call-by-reference. ${ }^{8}$

Here are examples:

```
MODULE main
    ..
    VAR
        a : boolean;
        b : foo(a);
MODULE foo(x)
    ASSIGN
        x := TRUE;
```

the variable $a$ is assigned the value TRUE. This distinguishes the call-by-reference mechanism from a call-by-value scheme.
Now consider the following program:

```
MODULE main
    DEFINE
        a := 0;
    VAR
        b : bar(a);
.
MODULE bar(x)
    DEFINE
        a := 1;
        y := x;
```

In this program, the value of y is 0 . On the other hand, using a call-by-name mechanism, the value of y would be 1 , since a would be substituted as an expression for x .
Forward references to module names are allowed, but circular references are not, and result in an error.
Note: NUXMV does no longer support the keyword process.

[^5]
### 2.3.12 References to Module Components (Variables and Defines)

As described in Section 2.2 .3 [Variables and Defines], page 14, defines and variables can be referenced in expressions as variable_identifiers and define_identifiers respectively, both of which are complex identifiers. The syntax of a complex identifier is:

```
complex_identifier ::
    identifier
    | complex_identifier . identifier
    complex_identifier [ simple_expression ]
    self
```

Every variable and define used in an expression should be declared. It is possible to have forward references when a variable or define identifier is used textually before the corresponding declaration.

Notations with . (<DOT>) are used to access the components of modules. For example, if $m$ is an instance of a module (see Section 2.3.11 [MODULE Instantiations], page 31 for information about instances of modules) then the expression $\mathrm{m} . \mathrm{c}$ identifies the component c of the module instance m . This is precisely analogous to accessing a component of a structured data type.

Note that actual parameters of a module can potentially be instances of other modules. Therefore, parameters of modules allow access to the components of other module instances, as in the following example:

```
MODULE main
... VAR
    a : bar;
    m : foo(a);
...
MODULE bar
    VAR
        q : boolean;
        p : boolean;
MODULE foo(c)
    DEFINE
        flag := c.q | c.p;
```

Here, the value of 'm.flag' is the logical OR of 'a.p' and 'a.q'.
Individual elements of an array are accessed in the typical fashion with the index given in square brackets. See 2.2.3 for more information.

It is possible to refer to the name that the current module has been instantiated to by using the self built-in identifier.

```
MODULE container(init_value1, init_value2)
    VAR c1 : counter(init_value1, self);
    VAR c2 : counter(init_value2, self);
MODULE counter(init_value, my_container)
    VAR v: 1..100;
    ASSIGN
        init(v) := init_value;
    DEFINE
        greatestCounterInContainer := v >= my_container.c1.v &
                        v >= my_container.c2.v;
MODULE main
    VAR c : container(14, 7);
    SPEC
        c.c1.greatestCounterInContainer;
```

In this example an instance of the module container is passed to the sub-module counter. In the main module, c is declared to be an instance of the module container, which declares two instances of the module counter. Every instance of the counter module has a define greatestCounterInContainer which specifies the condition when this particular counter has the greatest value in the container it belongs to. So a counter needs access to the parent container to access all the counters in the container.

### 2.3.13 A Program and the main Module

The syntax of a NUXMV program is:

```
program :: module_list
module_list ::
    module
    | module_list module
```

There must be one module with the name main and no formal parameters. The module main is the one evaluated by the interpreter.

### 2.3.14 Namespaces and Constraints on Declarations

Identifiers in the NUXMV input language may reference five different entities: modules, variables, defines, module instances, and symbolic constants.

Module identifiers have their own separate namespace. Module identifiers can be used in module type specifiers only, and no other kind of identifiers can be used there (see Section 2.3.11 [MODULE Instantiations], page 31). Thus, module identifiers may be equal to other kinds of identifiers without making the program ambiguous. However, no two modules should be declared with the same identifier. Modules cannot be declared in other modules, therefore they are always referenced by simple identifiers.

Variable, define, and module instance identifiers are introduced in a program when the module containing their declarations is instantiated. Inside this module the variables, defines and module instances may be referenced by the simple identifiers. Inside other modules, their simple identifiers should be preceded by the identifier of the module instance containing their declaration and ( $<D O T\rangle$ ). Such identifiers are called complex identifier. The full identifier is a complex identifier which references a variable, define, or a module instance from inside the main module.

Let us consider the following:

```
MODULE main
    VAR a : boolean;
    VAR b : foo;
    VAR c : moo;
MODULE foo
    VAR q : boolean;
        e : moo;
MODULE moo
    DEFINE f := 0 < 1;
MODULE not_used
    VAR n : boolean;
    VAR t : used;
MODULE used
    VAR k : boolean;
```

The full identifier of the variable $a$ is $a$, the full identifier of the variable $q$ (from the module foo) is $b . q$, the full identifier of the module instance $e$ (from the module foo) is $b . e$, the full identifiers of the define $f$ (from the module moo) are b.e.f and c.f, because two module instances contain this define. Notice that, the variables $n$
and $k$ as well as the module instance $t$ do not have full identifiers because they cannot be accessed from main (since the module not used is not instantiated).

In the nUXmv language, variable, define, and module instances belong to one namespace, and no two full identifiers of different variable, define, or module instances should be equal. Also, none of them can be redefined.

A symbolic constant can be introduced by a variable declaration if its type specifier enumerates the symbolic constant. For example, the variable declaration

VAR a : \{OK, FAIL, waiting\};
declares the variable a as well as the symbolic constants OK, FAIL and waiting. The full identifiers of the symbolic constants are equal to their simple identifiers with the additional condition - the variable whose declaration declares the symbolic constants also has a full identifier.

Symbolic constants have a separate namespace, so their identifiers may potentially be equal, for example, variable identifiers. It is an error, if the same identifier in an expression can simultaneously refer to a symbolic constant and a variable or a define. A symbolic constant may be declared an arbitrary number of times, but it must be declared at least once, if it is used in an expression.

### 2.3.15 Context

Every module instance has its own context, in which all expressions are analyzed. The context can be defined as the full identifiers of variables declared in the module without their simple identifiers. Let us consider the following example:

```
MODULE main
    VAR a : foo;
    VAR b : moo;
MODULE foo
    VAR c : moo;
MODULE moo
    VAR d : boolean;
```

The context of the module main is ' ${ }^{\prime}(\mathrm{empty})^{9}$, the context of the module instance a (and inside the module foo) is 'a.', the contexts of module moo may be 'b.' (if the module instance b is analyzed) and 'a.c.' (if the module instance a.c is analyzed).

### 2.3.16 ISA Declarations

There are cases in which some parts of a module could be shared among different modules, or could be used as a module themselves. Similarly to NuSMV, in NUXmV it is possible to declare the common parts as separate modules, and then use the ISA declaration to import the common parts inside a module declaration. The syntax of an isa_declaration is as follows:

```
isa_declaration :: ISA identifier
```

where identifier must be the name of a declared module. The ISA_declaration can be thought as a simple macro expansion command, because the body of the module referenced by an ISA command is replaced to the ISA_declaration.

Warning: ISA is a deprecated feature and will be removed from future versions of NUXMV. Therefore, avoid the use of ISA_declarations. Use module instances instead.

### 2.3.17 PRED and MIRROR Declarations

When using abstraction-based techniques, such as Counterexample Guided Abstraction Refinement [CGJ ${ }^{+}$03] or k -induction with implicit abstraction [Ton09], one may want to declare an initial set of predicates to be used in the model. This is possible by using the keyword Pred.

The syntax of a pred_declaration is as follows:

[^6]```
pred_declaration :: PRED simple_expression [;]
    PRED < identifier > := simple_expression [;]
```

where identifier is an arbitrary name to be assigned to the predicate.
Another way to specify the predicates to be used in the abstraction-based techniques model consists in "mirroring", i.e. preserving in the abstract space, a variable. A mirrored variable with its type will be declared in the abstract model as it was declated in the concrete model. Mirrored variables are introduced with the keyword MIRROR. A mirror_declaration is as follows.

```
mirror_declaration :: MIRROR variable_identifier [;]
```


### 2.4 Specifications

The specifications to be checked on the FSM can be expressed in temporal logics like Computation Tree Logic (CTL), Linear Temporal Logic (LTL)extended with Past Operators, and Property Specification Language (PSL) [psl03] that includes CTL and LTL with Sequential Extended Regular Expressions (SERE), a variant of classical regular expressions. It is also possible to analyze quantitative characteristics of the FSM by specifying real-time CTL specifications. Specifications can be positioned within modules, in which case they are preprocessed to rename the variables according to their context.

CTL and LTL specifications are evaluated by nUXmV in order to determine their truth or falsity in the FSM. When a specification is discovered to be false, NUXMV constructs and prints a counterexample, i.e. a trace of the FSM that falsifies the property.

### 2.4.1 CTL Specifications

A CTL specification is given as a formula in the temporal logic CTL, introduced by the keyword 'CTLSPEC' (however, deprecated keyword 'SPEC' can be used instead.) The syntax of this specification is:

```
ctl_specification :: CTLSPEC ctl_expr [;]
    SPEC ctl_expr [;]
CTLSPEC NAME name := ctl_expr [;]
SPEC NAME name := ctl_expr [;]
```

The syntax of CTL formulas recognized by NUXMV is as follows:

```
ctl_expr ::
    simple_expr -- a simple boolean expression
    | ( ctl_expr )
    | ! ctl_expr -- logical not
    | ctl_expr & ctl_expr -- logical and
    | ctl_expr | ctl_expr -- logical or
    | ctl_expr xor ctl_expr -- logical exclusive or
    | ctl_expr xnor ctl_expr -- logical NOT exclusive or
    | ctl_expr -> ctl_expr -- logical implies
    | ctl_expr <-> ctl_expr -- logical equivalence
    | EG ctl_expr -- exists globally
    | EX ctl_expr -- exists next state
    | EF ctl_expr -- exists finally
    | AG ctl_expr -- forall globally
    AX ctl_expr -- forall next state
    | AF ctl_expr -- forall finally
    E [ ctl_expr U ctl_expr ] -- exists until
    A [ ctl_expr U ctl_expr ] -- forall until
```

Since simple_expr cannot contain the next operator, ctl_expr cannot contain it either. The ctl_expr should also be a boolean expression.

Intuitively the semantics of CTL operators is as follows:


- EX $p$ is true in a state $s$ if there exists a state $s^{\prime}$ such that a transition goes from $s$ to $s^{\prime}$ and $p$ is true in $s^{\prime}$.
- AX $p$ is true in a state $s$ if for all states $s^{\prime}$ where there is a transition from $s$ to $s^{\prime}, p$ is true in $s^{\prime}$.
- EF $p$ is true in a state $s_{0}$ if there exists a series of transitions $s_{0} \rightarrow s_{1}, s_{1} \rightarrow s_{2}, \ldots, s_{n-1} \rightarrow s_{n}$ such that $p$ is true in $s_{n}$.
- AF $p$ is true in a state $s_{0}$ if for all series of transitions $s_{0} \rightarrow s_{1}, s_{1} \rightarrow s_{2}, \ldots, s_{n-1} \rightarrow s_{n} p$ is true in $s_{n}$.
- EG $p$ is true in a state $s_{0}$ if there exists an infinite series of transitions $s_{0} \rightarrow s_{1}, s_{1} \rightarrow s_{2}, \ldots$ such that $p$ is true in every $s_{i}$.
- AG $p$ is true in a state $s_{0}$ if for all infinite series of transitions $s_{0} \rightarrow s_{1}, s_{1} \rightarrow s_{2}, \ldots p$ is true in every $s_{i}$.
- $\mathbf{E}[p \mathrm{U} q]$ is true in a state $s_{0}$ if there exists a series of transitions $s_{0} \rightarrow s_{1}, s_{1} \rightarrow s_{2}, \ldots, s_{n-1} \rightarrow s_{n}$ such that $p$ is true in every state from $s_{0}$ to $s_{n-1}$ and $q$ is true in state $s_{n}$.
- A[p $\operatorname{U} q]$ is true in a state $s_{0}$ if for all series of transitions $s_{0} \rightarrow s_{1}, s_{1} \rightarrow s_{2}, \ldots, s_{n-1} \rightarrow s_{n} p$ is true in every state from $s_{0}$ to $s_{n-1}$ and $q$ is true in state $s_{n}$.

A CTL formula is true if it is true in all initial states.
For a detailed description about the semantics of PSL operators, please see [ps103].

### 2.4.2 Invariant Specifications

It is also possible to specify invariant specifications with special constructs. Invariants are propositional formulas which must hold invariantly in the model. The corresponding command is INVARSPEC, with syntax:

```
invar_specification :: INVARSPEC next_expr ;
    INVARSPEC NAME name := next_expr [;]
```

This statement is intuitively equivalent to

```
CTLSPEC AG simple_expr ;
```

but can be checked by a specialised algorithm during reachability analysis and Invariant Specifications can contain next operators. Fairness constraints are not taken into account during invariant checking.

### 2.4.3 LTL Specifications

LTL specifications are introduced by the keyword LTLSPEC. The syntax of this specification is:

```
ltl_specification :: LTLSPEC ltl_expr [;]
    LTLSPEC NAME name := ltl_expr [;]
```

The syntax of LTL formulas recognized by NUXMV is as follows:

```
ltl_expr ::
    next_expr -- a next boolean expression
    | ( ltl_expr )
    | ! ltl_expr -- logical not
    | ltl_expr & ltl_expr -- logical and
    | ltl_expr | ltl_expr -- logical or
    | ltl_expr xor ltl_expr -- logical exclusive or
    | ltl_expr xnor ltl_expr -- logical NOT exclusive or
    | ltl_expr -> ltl_expr -- logical implies
    | ltl_expr <-> ltl_expr -- logical equivalence
    -- FUTURE
    | X ltl_expr -- next state
    | G ltl_expr -- globally
    | F ltl_expr -- finally
    | ltl_expr U ltl_expr -- until
```

| $\begin{aligned} & \text { \| ltl_expr v ltl_expr } \\ & \text {-- PAST } \end{aligned}$ | -- releases |
| :---: | :---: |
| \| Y ltl_expr | -- previous state |
| \| $\mathbf{Z}$ ltl_expr | -- not previous state not |
| \| H ltl_expr | -- historically |
| \| O ltl_expr | once |
| \| ltl_expr S ltl_expr | - since |
| \| ltl_expr T ltl_expr | -- triggered |

Intuitively the semantics of LTL operators is as follows:

- $\mathbf{X} p$ is true at time $t$ if $p$ is true at time $t+1$.
- $\mathbf{F} p$ is true at time $t$ if $p$ is true at some time $t^{\prime} \geq t$.
- G $p$ is true at time $t$ if $p$ is true at all times $t^{\prime} \geq t$.
- $p \mathrm{U} q$ is true at time $t$ if $q$ is true at some time $t^{\prime} \geq t$, and for all time $t^{\prime \prime}$ (such that $t \leq t^{\prime \prime}<t^{\prime}$ ) $p$ is true.
- $p \mathbf{v} q$ is true at time $t$ if $q$ holds at all time steps $t^{\prime} \geq t$ up to and including the time step $t^{\prime \prime}$ where $p$ also holds. Alternatively, it may be the case that $p$ never holds in which case $q$ must hold in all time steps $t^{\prime} \geq t$.
- $\mathbf{Y} p$ is true at time $t>t_{0}$ if $p$ holds at time $t-1$. $\mathbf{Y} p$ is false at time $t_{0}$.
- $\mathbf{z} p$ is equivalent to $\mathbf{Y} p$ with the exception that the expression is true at time $t_{0}$.
- н $p$ is true at time $t$ if $p$ holds in all previous time steps $t^{\prime} \leq t$.
- $0 p$ is true at time $t$ if $p$ held in at least one of the previous time steps $t^{\prime} \leq t$.
- p $\mathbf{S} q$ is true at time $t$ if $q$ held at time $t^{\prime} \leq t$ and $p$ holds in all time steps $t^{\prime \prime}$ such that $t^{\prime}<t^{\prime \prime} \leq t$.
- $p \mathbf{T} q$ is true at time $t$ if $p$ held at time $t^{\prime} \leq t$ and $q$ holds in all time steps $t^{\prime \prime}$ such that $t^{\prime} \leq t^{\prime \prime} \leq t$. Alternatively, if $p$ has never been true, then $q$ must hold in all time steps $t^{\prime \prime}$ such that $t_{0} \leq t^{\prime \prime} \leq t$

An LTL formula is true if it is true at the initial time $t_{0}$.
In NUXMV, LTL specifications can be analyzed, depending on the fact that the model is finite-state or infinite state, by means of BDD-based reasoning, by means of SAT-based techniques or for SMT based techniques. In the case of BDD-based reasoning, nUXMV proceeds according to [CGH97a]. For each LTL specification, a tableau of the behaviors falsifying the property is constructed, and then synchronously composed with the model. Similarly to NUSMV, the [CGH97a] approach is fully integrated within NUXMV, and allows full treatment of past temporal operators. Note that the counterexample is generated in such a way to show that the falsity of a LTL specification may contain state variables which have been introduced by the tableau construction procedure.

In the case of SAT/SMT-based reasoning, a similar tableau construction is carried out to encode the paths of limited length, violating the property. NUXMV, similarly to NUSMV, generates a propositional satisfiability problem, that is then tackled by means of efficient SAT or SMT solvers.

In all cases, the tableau constructions are completely transparent to the user.

## Important Difference Between BDD and SAT/SMT Based LTL Model Checking

If a FSM to be checked is not total (i.e. it has at least a deadlock state) the model checking may return different results for the same LTL specification depending on the verification engine used. For example, let us consider the model below.

```
MODULE main
VAR s : boolean;
TRANS s = TRUE
LTLSPEC G (s = TRUE)
```

The LTL specification is proved valid by BDD－based model checking but is violated by SAT／SMT－based bounded model checking．The counter－example found consists of one state s＝FALSE．

This difference between the results is caused by the fact that BDD model checking investigates only infinite paths whereas SAT／SMT－based model checking is able to deal also with finite paths．Apparently infinite paths cannot ever have $s=F A L S E$ as then the transition relation will not hold between the consecutive states in the path． A finite path consisting of just one state $s=F A L S E$ violates the specification $G(s=$ TRUE $)$ and is still consistent with the FSM as the transition relation is not taken ever and there is not initial condition to violate．Note however that this state is a deadlock and cannot have consecutive states．

In order to make SAT／SMT－based bound model checking ignore finite paths it is enough to add a fairness condition to the main module：

## JUSTICE TRUE；

Being limited to fair paths，SAT／SMT－based bounded model checking cannot find a finite counter－example and results of model checking become consistent with BDD－based model checking．

## 2．4．4 Real Time CTL Specifications and Computations

nuXmV（as well as NuSMV）allows for Real Time CTL specifications［EMSS91］．Similarly to NUSMV，in nUXMV we assume that each transition takes unit time for execution．RTCTL extends the syntax of CTL path expressions with the following bounded modalities：

```
rtctl_expr ::
        ctl_expr
        EBF range rtctl_expr
        ABF range rtctl_expr
        EBG range rtctl_expr
        ABG range rtctl_expr
        A [ rtctl_expr BU range rtctl_expr ]
        E [ rtctl_expr BU range rtctl_expr ]
range :: integer_number .. integer_number
```

Given ranges must be non－negative．
Intuitively，the semantics of the RTCTL operators is as follows：
－EBF m．．n $p$ requires that there exists a path starting from a state，such that property $p$ holds in a future time instant $i$ ，with $m \leq i \leq n$
－ABF m．．n $p$ requires that for all paths starting from a state，property $p$ holds in a future time instant $i$ ，with $m \leq i \leq n$
－EBG $m \ldots n$ requires that there exists a path starting from a state，such that property $p$ holds in all future time instants $i$ ，with $m \leq i \leq n$
－ABG m．．n $p$ requires that for all paths starting from a state，property $p$ holds in all future time instants $i$ ， with $m \leq i \leq n$
－E［ $p \mathbf{B U} m . . n q$ ］requires that there exists a path starting from a state，such that property $q$ holds in a future time instant $i$ ，with $m \leq i \leq n$ ，and property $p$ holds in all future time instants $j$ ，with $m \leq j<i$
－A［ $p$ BU m．．n $q$ ］，requires that for all paths starting from a state，property $q$ holds in a future time instant $i$ ，with $m \leq i \leq n$ ，and property $p$ holds in all future time instants $j$ ，with $m \leq j<i$

Real time CTL specifications can be defined with the following syntax，which extends the syntax for CTL speci－ fications．（keyword＇SPEC＇is deprecated）

```
rtctl_specification :: CTLSPEC rtctl_expr [;]
    | SPEC rtctl_expr [;]
    | CTLSPEC NAME name := rtctl_expr [;]
    | SPEC NAME name := rtctl_expr [;]
```

With the COMPUTE statement, it is also possible to compute quantitative information on the FSM. In particular, it is possible to compute the exact bound on the delay between two specified events, expressed as CTL formulas. The syntax is the following:

```
compute_specification :: COMPUTE compute_expr [;]
    COMPUTE NAME name := compute_expr [;]
```

where

```
compute_expr :: MIN [ rtctl_expr , rtctl_expr ]
```

    MAX [ rtctl_expr , rtctl_expr ]
    MIN [start, final] returns the length of the shortest path from a state in start to a state in final. For this, the set of states reachable from start is computed. If at any point, we encounter a state satisfying final, we return the number of steps taken to reach the state. If a fixed point is reached and no computed states intersect final then infinity is returned.
MAX [start, final] returns the length of the longest path from a state in start to a state in final. If there exists an infinite path beginning in a state in start that never reaches a state in final, then infinity is returned. If any of the initial or final states is empty, then undefined is returned.

It is important to remark here that if the FSM is not total (i.e. it contains deadlock states) COMPUTE may produce wrong results. It is possible to check the FSM against deadlock states by calling the command check_f.sm.

### 2.4.5 PSL Specifications

NUXMV, similarly to NUSMV, allows to specify PSL properties that comply with version 1.01 of PSL Language Reference Manual [psl03]. PSL specifications are introduced by the keyword "PSLSPEC". The syntax of this declaration (as from the PSL parsers distributed by IBM, [PSL]) is:

```
pslspec_declaration :: PSLSPEC psl_expr [;]
PSLSPEC NAME name := psl_expr [;]
```

where

```
psl_expr ::
    psl_primary_expr
    psl_unary_expr
    psl_binary_expr
    psl_conditional_expr
    psl_case_expr
    psl_property
```

The first five classes define the building blocks for psl_property and provide means of combining instances of that class; they are defined as follows:

```
psl_primary_expr ::
    number ;; a numeric constant
    boolean ;; a boolean constant
    word ;; a word constant
    var_id ;; a variable identifier
    { psl_expr , ... , psl_expr }
    { psl_expr "{" psl_expr , ... , "psl_expr" }}
    ( psl_expr )
psl_unary_expr ::
    + psl_primary_expr
    - psl_primary_expr
    ! psl_primary_expr
    bool ( psl_expr )
    word1 ( psl_expr )
    uwconst ( psl_expr, psl_expr )
```

```
    swconst ( psl_expr, psl_expr )
    sizeof ( psl_expr )
    toint ( psl_expr )
    signed ( psl_expr )
    unsigned ( psl_expr )
    extend ( psl_expr, psl_primary_expr )
    resize ( psl_expr, psl_primary_expr )
    select ( psl_expr, psl_expr, psl_expr )
psl_binary_expr ::
    psl_expr + psl_expr
    psl_expr union psl_expr
    psl_expr in psl_expr
    psl_expr - psl_expr
    psl_expr * psl_expr
    psl_expr / psl_expr
    psl_expr % psl_expr
    psl_expr == psl_expr
    psl_expr != psl_expr
    psl_expr < psl_expr
    psl_expr <= psl_expr
    psl_expr > psl_expr
    psl_expr >= psl_expr
    psl_expr & psl_expr
    psl_expr | psl_expr
    psl_expr xor psl_expr
    psl_expr xnor psl_expr
    psl_expr << psl_expr
    psl_expr >> psl_expr
    psl_expr :: psl_expr
psl_conditional_expr ::
    psl_expr ? psl_expr : psl_expr
psl_case_expr ::
case
    psl_expr : psl_expr ;
    psl_expr : psl_expr ;
endcase
```

Among the subclasses of psl_expr we depict the class psl_bexpr that will be used in the following to identify purely boolean, i.e. not temporal, expressions. The class of PSL properties psl_property is defined as follows:

```
psl_property ::
    replicator psl_expr ; ; a replicated property
    FL_property abort psl_bexpr
    psl_expr <-> psl_expr
    psl_expr -> psl_expr
    FL_property
    OBE_property
replicator ::
    forall var_id [index_range] in value_set :
index_range ::
    [ range ]
range ::
    low_bound : high_bound
low_bound ::
    number
    | identifier
high_bound ::
    number
```

```
    identifier
    inf ;; inifite high bound
value_set ::
    { value_range , ... , value_range }
    boolean
value_range ::
    psl_expr
    range
```

The instances of FL_property are temporal properties built using LTL operators and SEREs operators, and are defined as follows:

```
FL_property ::
    ;; PRIMITIVE LTL OPERATORS
        X FL_property
    X! FL_property
    F FL_property
    G FL_property
    [ FL_property U FL_property ]
    [ FL_property W FL_property ]
    ; SIMPLE TEMPORAL OPERATORS
    always FL_property
    never FL_property
    next FL_property
    next! FL_property
    eventually! FL_property
    FL_property until! FL_property
    FL_property until FL_property
    FL_property until!_ FL_property
    FL_property until_ FL_property
    FL_property before! FL_property
    FL_property before FL_property
    FL_property before!_ FL_property
    FL_property before_ FL_property
    ; EXTENDED NEXT OPERATORS
    X [number] ( FL_property )
    X! [number] ( FL_property )
    next [number] ( FL_property )
    next! [number] ( FL_property )
    ;
    next_a [range] ( FL_property )
    next_a! [range] ( FL_property )
    next_e [range] ( FL_property )
    next_e! [range] ( FL_property )
;;
    next_event! ( psl_bexpr ) ( FL_property )
    next_event ( psl_bexpr ) ( FL_property )
    next_event! ( psl_bexpr ) [ number ] ( FL_property )
    next_event ( psl_bexpr ) [ number ] ( FL_property )
;;
    next_event_a! ( psl_bexpr ) [psl_expr] ( FL_property )
    next_event_a ( psl_bexpr ) [psl_expr] ( FL_property )
    next_event_e! ( psl_bexpr ) [psl_expr] ( FL_property )
    next_event_e ( psl_bexpr ) [psl_expr] ( FL__property )
    OPERATORS ON SERES
    sequence ( FL_property )
    sequence |-> sequence [!]
    sequence |=> sequence [!]
;;
    always sequence
```

$\underset{\substack{\text { FoNDOAIONE } \\ \text { BRUNO KESSLER }}}{ }$

```
G sequence
never sequence
eventually! sequence
;;
within! ( sequence_or_psl_bexpr , psl_bexpr ) sequence
within ( sequence_or_psl_bexpr , psl_bexpr ) sequence
within!_ ( sequence_or_psl_bexpr , psl_bexpr ) sequence
within_ ( sequence_or_psl_bexpr , psl_bexpr ) sequence
;
whilenot! ( psl_bexpr ) sequence
whilenot ( psl_bexpr ) sequence
whilenot!_ ( psl_bexpr ) sequence
whilenot_ ( psl_bexpr ) sequence
sequence_or_psl_bexpr ::
    sequence
psl_bexpr
```

Sequences, i.e. istances of class sequence, are defined as follows:

```
sequence ::
    { SERE }
SERE ::
    sequence
    psl_bexpr
;; COMPOSITION OPERATORS
    SERE ; SERE
    SERE : SERE
    SERE & SERE
    SERE && SERE
    SERE | SERE
    ; RegExp QUALIFIERS
    SERE [* [count] ]
    [* [count] ]
    SERE [+]
    [+]
;;
    psl_bexpr [= count ]
    psl_bexpr [-> count ]
count ::
    number
    range
```

Istances of OBE_property are CTL properties in the PSL style and are defined as follows:

```
OBE_property ::
    AX OBE_property
    AG OBE_property
    AF OBE_property
    A [ OBE_property U OBE_property ]
    EX OBE_property
    EG OBE_property
    EF OBE_property
    E [ OBE_property U OBE_property ]
```

The NUXMV parser allows to input any specification based on the grammar above, but currently, verification of PSL specifications is supported only for the OBE subset, and for a subset of PSL for which it is possible to define a translation into LTL. For the specifications that belong to these subsets, it is possible to apply all the verification techniques that can be applied to LTL and CTL Specifications.

Note: Full support for PSL will be integrated in forthcoming releases of NUXMV.

### 2.5 Variable Order Input

As it is the case in NUSMV, NUXMV allows to specify the order in which variables should appear in the generated BDDs. The file which gives the desired order can be read in using the -i option in batch mode or by setting the input_order_file environment variable in interactive mode. ${ }^{10}$

### 2.5.1 Input File Syntax

The syntax for input files describing the desired variable ordering is as follows, where the file can be considered as a list of variable names, each of which must be on a separate line:

```
vars_list :: EMPTY
    | var_list_item vars_list
var_list_item :: complex_identifier
    | complex_identifier . integer_number
```

Where EMPTY means parsing nothing.
This grammar allows for parsing a list of variable names of the following forms:

```
Complete_Var_Name -- to specify an ordinary variable
Complete_Var_Name[index] -- to specify an array variable element
Complete_Var_Name.NUMBER -- to specify a specific bit of a
    -- scalar variable
```

where Complete_Var_Name is just the name of the variable if it appears in the module MAIN, otherwise it has the module name(s) prepended to the start, for example:
mod1.mod2...modN.varname
where varname is a variable in modn, and modn. varname is a variable in modn-1, and so on. Note that the module name main is implicitely prepended to every variable name and therefore must not be included in their declarations.
Any variable which appears in the model file, but not the ordering file is placed after all the others in the ordering. Variables which appear in the ordering file but not the model file are ignored. Similarly to NUSMV, in both cases nUXMV displays a warning message stating these actions.

Comments can be included by using the same syntax as regular NuSMV files. That is, by starting the line with --.

### 2.5.2 Scalar Variables

A variable, which has a finite range of values that it can take, is encoded as a set of boolean variables (i.e. bits). These boolean variables represent the binary equivalents of all the possible values for the scalar variable. Thus, a scalar variable that can take values from 0 to 7 would require three boolean variables to represent it.

It is possible not only to declare the position of a scalar variable in the ordering file, but each of the boolean variables which represent it.
If only the scalar variable itself is named then all the boolean variables which are actually used to encode it are grouped together in the BDD package.
Variables which are grouped together will always remain next to each other in the BDD package and in the same order. When dynamic variable re-ordering is carried out, the group of variables are treated as one entity and moved as such.
If a scalar variable is omitted from the ordering file then it will be added at the end of the variable order and the specific-bit variables that represent it will be grouped together. However, if any specific-bit variables have been declared in the ordering file (see below) then these will not be grouped with the remaining ones.

[^7]It is also possible to specify the location of specific bit variables anywhere in the ordering. This is achieved by first specifying the scalar variable name in the desired location, then simply specifying Complete_Var_Name.i at the position where you want that bit variable to appear:

```
Complete_Var_Name
Complete_Var_Name.i
```

The result of doing this is that the variable representing the $i^{t h}$ bit is located in a different position to the remainder of the variables representing the rest of the bits. The specific-bit variables varname. $0, \ldots$, varname.i-1, varname. $i+1, \ldots$, varname. $N$ are grouped together as before.

If any one bit occurs before the variable it belongs to, the remaining specific-bit variables are not grouped together:

```
Complete_Var_Name.i
Complete_Var_Name
```

The variable representing the $i^{t h}$ bit is located at the position given in the variable ordering and the remainder are located where the scalar variable name is declared. In this case, the remaining bit variables will not be grouped together.
This is just a short-hand way of writing each individual specific-bit variable in the ordering file. The following are equivalent:

```
Complete_Var_Name.0 Complete_Var_Name.0
Complete_Var_Name.1 Complete_Var_Name
\vdots
Complete_Var_Name.N-1
```

where the scalar variable Complete_Var_Name requires N boolean variables to encode all the possible values that it may take. It is still possible to then specify other specific-bit variables at later points in the ordering file as before.

### 2.5.3 Array Variables

When declaring array variables in the ordering file, each individual element must be specified separately. It is not permitted to specify just the name of the array. The reason for this is that the actual definition of an array in the model file is essentially a shorthand method of defining a list of variables that all have the same type. Nothing is gained by declaring it as an array over declaring each of the elements individually, and there is no difference in terms of the internal representation of the variables.

### 2.6 Clusters Ordering

When nUXMV builds a clusterized BDD-based FSM during model construction (as it is the case for NUSMV), an initial simple clusters list is roughly constructed by iterating through a list of variables, and by constructing the clusters by picking the transition relation associated to each variable in the list. Later, the clusters list will be refined and improved by applying the clustering alghorithm that the user previoulsy selected (for further information, see partitioning methods in the NUSMV user manual $\left.\left[\mathrm{CCCJ}^{+} 10\right]\right)$.

NUXMV, similar to NUSMV, allows to specify an ordering for the initial list of variables that is used to build the clusters. The option trans_order_file can be used to specify a file containing a variable ordering. This feature is inherited directly from NUSMV (for further information see the NUSMV user manual [CCCJ ${ }^{+}$10]).

Grammar of the clusters ordering file in NUXMV is the same of the one in the NUSMV user manual $\left[\mathrm{CCCJ}^{+} 10\right]$.

## Chapter 3

## Running nUXmV interactively

nuXMV inherits from NuSMV the interactive shell. In this mode nUXMV, like NUSMV, enters a read-eval-print loop. The user can activate the various NUXMV computation steps as system commands with different options. These steps can therefore be invoked separately, possibly undone or repeated under different modalities. As for NUSMV, the interactive shell of NUXMV is activated from the system prompt as follows ('nuXmv >' is the default nUXMV shell prompt):

```
system_prompt> nuXmv -int <RET>
nuXmv >
```

When running interactively, NUXMV first tries to read and execute commands from an initialization file if such file can be found and is readable unless -s is passed on the command line.
Search is done in this order:

1. File master.nuxmvrc is looked for in directory defined in environment variable NUXMV_LIBRARY_PATH or in default library path (e.g. /usr/local/share/nusmv under GNU/Linux) if no such variable is defined.
2. If no such file exists, file . nuxmvrc is looked for in user's home directory.
3. If no such file exists, . nuxmvrc is looked for in current directory.

Tip: To see which file are searched for and the search paths we recommend to use the command line option -h and to look at the description for option -s.

Commands in the initialization file (if any) are executed consecutively. When initialization phase is completed the nUXMV shell is displayed and the system is now ready to execute user commands.

Similar to NuSMV, a nUXmV command is a sequence of words. The first word specifies the command to be executed. The remaining words are arguments to the invoked command. Commands separated by a ';' are executed sequentially; the NUXMV shell waits for each command to terminate in turn. The behavior of commands can depend on environment variables, similar to "csh" environment variables.

It is also possible to make NUXMV read and execute a sequence of commands from a file, through the command line option -source:

```
system_prompt> nuXmv -source cmd_file <RET>
```

-source cmd-file Starts the interactive shell and then executes NUXMV commands from file cmd-file. If an error occurs during a command execution, commands that follow will not be executed. See also the variable on_failure_script_quits. The option-source implies -int.

Command sequences in NUXMV (similar to NUSMV) must obey the (partial) order specified in the Figure 3.1 depicted at page 47. For instance, it is not possible to evaluate CTL/LTL or invariant properties before the model is built. In Figure 3.1 a star $(*)$ is used as prefix to denote new commands provided by NUXMV, and not found in NUSMV. Furthermore, loop backs are not represented to ease the reading. Nevertheless, the flow should be still well comprehensible.

A number of commands and environment variables, like those dealing with file names, accept arbitrary strings. There are a few reserved characters which must be escaped if they are to be used literally in such situations. See the section describing the history command, on page 99, for more information.

The verbosity of NUXMV is controlled by the verbose_level environment variable.

## verbose level

Environment Variable
Controls the verbosity of the system. Possible values are integers from 0 (no messages) to 4 (full messages). The default value is 0 .
nUXMV provides to the user all the commands that NuSMV provides. Moreover, it extends the set of commands with new ones. In Chapter 4 we describe all the NUSMV commands, while in Chapter 5 we describe all the new NUXMV commands. All the commands are organizez for functionality.

## Command Annotations

All the NUSMV commands operates over finite domains. In the following we annotated each new commands with a flag stating whether the command is applicable only to finite-state domain, or if it can be applied to infinite-state domains (which include finite domains). We use the folloiwng annotations:

- [F]: the command can be applied only to finite-state domains.
- [I]: the command can be applied only to infinite-state domains, which include finite-state ones.
- [F,I]: the command provides the user with command line options to invoke the finite-state or the infinite-state version (that uses SMT) for the underlying algorithm.
$\underset{\substack{\text { Fonvazione } \\ \text { BRINOK KESS ER }}}{ }$


Figure 3.1: The dependency among nUXMV commands.

## Chapter 4

## Commands from NuSMV

In the following we present the commands inherited from NuSMV. We also describe the environment variables that may affect the behavior of the commands. All the commands have been classified in different categories.

### 4.1 Model Reading and Building

The following commands allow for the parsing and compilation of the model in order to enable all the verification algorithms.

```
read_model-Reads a NuSMV file into NuSMV.
```

```
read_model [-h] [-i model-file]
```

Reads a NUXMV file. If the -i option is not specified, it reads from the file specified in the environment variable input_file.

Command Options:
-i model-file Sets the environment variable input_file to model-file, and reads the model from the specified file.

## input file

Environment Variable
Stores the name of the input file containing the model. It can be set by the "set" command or by the command line option ' $-i$ '. There is no default value.

Stores the list of pre-processors to be run on the input file before it is parsed by nUXMV. The pre-processors are executed in the order specified by this variable. The argument must either be the empty string (specifying that no pre-processors are to be run on the input file), one single pre-processor name or a space seperated list of pre-processor names inside double quotes. Any invalid names are ignored. The default is none.

```
flatten_hierarchy [-h] [-d]
```

This command is responsible of the instantiation of modules and processes. The instantiation is performed by substituting the actual parameters for the formal parameters, and then by prefixing the result via the instance name.

Command Options:
-d Delays the construction of vars constraints until needed

## disable_syntactic_checks

Environment Variable
Enables or disables the syntactic checks that are performed by the "flatten_hierarchy" command. Warning: If the model is not well-formed, NUXMV may result in unpredictable results, use this option at your own risk.

## keep_single_value_vars

Environment Variable
Enables or disables the conversion of variables that can assume only one single possible value into constant DEFINEs.

## backward_compatibility <br> Environment Variable

As in NUSMV, this variable enables/disables type checking and other features provided by NUXMV. If set to 1 then the type checking is turned off, and nUXMV behaves as the old versions of NUSMV (for the pure boolean case) w.r.t. type checking and other features like writing of flattened and booleanized SMV files and promotion of boolean constants to their integer counterpart. If set to 0 then the type checking is turned on, and whenever a type error is encountered while compiling a NUXMV program the user is informed and the execution is stopped.

Since NUSMV 2.5.1, backward compatibility mode introduces a porting feature from old models which use constant 1 as case conditions, instead of forcing the use of TRUE.

The option by default it set to 0 .

## type_checking_warning_on

Environment Variable
Enables notification of warning messages generated by the type checking. If set to 0 , then messages are disregarded, otherwise if set to 1 they are notified to the user. As default it is set to 1 .

```
show_vars [-h] [-s] [-f] [-i] [-t | -V] [-v] [-m | -o output-file]
```

Prints a summary of the variables declared in the input file. Moreover, it prints also the list of symbolic input, frozen and state variables of the model with their range of values (as defined in the input file) if the proper command option is specified.

Command Options:

| $-s$ | Prints only state variables. |
| :--- | :--- |
| -f | Prints only frozen variables. |
| -i | Prints only input variables. |
| -t | Prints only the number of variables (among selected kinds), grouped by type. <br> -V |
|  | This option is incompatible with -V. |
|  | Prints only the list of variables with their types (among selected kinds), with <br> no summary information. This option is incompatible with -t. |

```
-v Prints verbosely. Scalar variable's values are not truncated if too long for printing.
-m Pipes the output to the program specified by the PAGER shell variable if defined, else through the UNIX command "more".
-o output-file Writes the output generated by the command to output-file.
```

show_dependencies - Shows the dependencies for the given expression
Command
show_dependencies [-h] [-k bound] -e expression
Prints the set of variables that are in the dependency set of the given expression. If the bound is specified using the -k argument, then the computation of the dependencies is done until the bound has been reached. If not specified, the computation is performed until no new dependencies are found.

Command Options:

```
-h Shows the command usage
-k bound Sets the bound limit for the dependencies computation
-e expr The expression on which the dependencies are computed
```

encode_variables - Builds the BDD variables necessary to compile the model
Command into a $B D D$.

```
encode_variables [-h] [-i order-file]
```

Generates the boolean BDD variables and the ADD needed to encode propositionally the (symbolic) variables declared in the model. The variables are created as default in the order in which they appear in a depth first traversal of the hierarchy.
The input order file can be partial and can contain variables not declared in the model. Variables not declared in the model are simply discarded. Variables declared in the model which are not listed in the ordering input file will be created and appended at the end of the given ordering list, according to the default ordering.

## Command Options:

-i order-file Sets the environment variable input_order_file to order-file, and reads the variable ordering to be used from file order-file. This can be combined with the write_order command. The variable ordering is written to a file, which can be inspected and reordered by the user, and then read back in.

## input_order file

Environment Variable
Indicates the file name containing the variable ordering to be used in building the model by the 'encode_variables' command. A value for this variable can also be provided with command line option -i. There is no default value.

Changes the behaviour of the command write_order.
When this variable is set, write_order will dump the bits constituting the boolean encoding of each scalar variable, instead of the scalar variable itself. This helps to work at bits level in the variable ordering file. See the command write_order for further information. The default value is 1 .

```
write_order [-h] [-b] [(-o | -f) order-file]
```

Writes the current order of BDD variables in the file specified via the -0 option. If no option is specified the environment variable output_order_file will be considered. If the variable output_order_file is unset (or set to an empty value) then standard output will be used.

By default, the bits constituting the scalar variables encoding are not dumped. When a variable bit should be dumped, the scalar variable which the bit belongs to is dumped instead if not previously dumped. The result is a variable ordering containing only scalar and boolean model variables.

To dump single bits instead of the corresponding scalar variables, either the option -b can be specified, or the environment variable write_order_dumps_bits must be previously set.
When the boolean variable dumping is enabled, the single bits will occur within the resulting ordering file in the same position that they occur at BDD level.

Command Options:

```
-b Dumps bits of scalar variables instead of the single scalar variables. See also
the variable write_order_dumps_bits.
Sets the environment variable output_order_file to order-file
and then dumps the ordering list into that file.
-f order-file
Alias for the -o option. Supplied for backward compatibility.
```


## output_order_file

Environment Variable
The file where the current variable ordering has to be written. A value for this variable can also be provided with command line option $-\mathbf{o}$. The default value is 'temp. ord'.

## vars_order_type

Environment Variable
Controls the manner variables are ordered by default, when a variable ordering is not specified by a user and not computed statically by heuristics (see variables input_order_file on page 50 and bdd_static_order_heuristics on page 52).

The individual bits of variables may or may not be interleaved. When bits interleaving is not used then bits belonging to one variable are grouped together in the ordering. Otherwise, the bits interleaving is applied and all higher bits of all variables are ordered before all the lower bits, i.e. N-th bits of all variables go before ( $\mathrm{N}-1$ )th bits. The exception is boolean variables which are ordered before variables of any other type though boolean variables consist of only 0 -th bit.
The value of vars_order_type may be:

- inputs_before. Input variables are forced to be ordered before state and frozen variables (default). No bits interleaving is done.
- inputs_after. Input variables are forced to be ordered after state and frozen variables. No bits interleaving is done.
- topological. Input, state and frozen variables are ordered as they are declared in the input smv file. No bits interleaving is done.
- inputs_before_bi. Bits are interleaved and in every group of N -th bits input variables are forced to be ordered before state and frozen variables. This is the default value.
- inputs_after_bi. Bits are interleaved and in every group of N -th bits input variables are forced to be ordered after state and frozen variables.
- topological_bi. Bits are interleaved and in every group of N-th bits input, state and frozen variables are ordered as they are declared in the input smv file.
- lexicographic. This is deprecated value. topological has to be used instead.


## bdd static_order heuristics

Environment Variable
When a variable ordering is not specified (see variable input_order_file on page 50) NUXMV can try to guess a good ordering by analyzing the input model.
Possible values are:

- none No heuristics are applied.
- basic This heuristics creates some initial ordering and then moves scalar and word variables in this ordering to form groups. Groups go one after another and every group contains variables which interact with each other in the model. For example, having variables $a, b, c, d, e, f$ and a single model constraint TRANS next (a)=b+1 -> (next (c)=d/e \& next (f)!=a) will results in 2 groups of variables $\{a, b, f\}$ and $\{c, d, e\}$.
Shell variable vars_order_type (page 51) provides additional control over the heuristics. In particular, it allows to put input/state variables in the initial ordering at the begin, the end or in topological order. Moreover, if the value of this variable is ending in _bi then in very individual group the bits of variables are additionally interleaved.
Note that variable groups created by the heuristics has nothing to do with BDD package groups which disallow dynamic reordering of variables in one group. After the heuristics is applied the dynamic reordering may move any bit of any variable at any position.
build_model - Compiles the flattened hierarchy into a BDD
build_model [-h] [-f] [-m Method]
Compiles the flattened hierarchy into a BDD (initial states, invariants, and transition relation) using the method specified in the environment variable partition_method for building the transition relation.

Command Options:

```
-m Method
-f Forces model construction. By default, only one partition method is allowed.
    This option allows to overcome this default, and to build the transition rela-
    tion with different partitioning methods.
```

The method to be used in building the transition relation, and to compute images and preimages. Possible values are:

- Monolithic. No partitioning at all.
- Threshold. Conjunctive partitioning, with a simple threshold heuristic. Assignments are collected in a single cluster until its size grows over the value specified in the variable conj_part_threshold. It is possible (default) to use affinity clustering to improve model checking performance. See affinity variable.
- Iwls95CP. Conjunctive partitioning, with clusters generated and ordered according to the heuristic described in [RAP $\left.{ }^{+} 95\right]$. Works in conjunction with the variables image_cluster_size, image_W1, image_W2, image_W3, image_W4. It is possible (default) to use affinity clustering to improve model checking performance. See affinity variable. It is also possible to avoid (default) preordering of clusters (see $\left[\mathrm{RAP}^{+} 95\right]$ ) by setting the iwls 95 preorder variable appropriately.


## conj_part_threshold

Environment Variable
The limit of the size of clusters (expressed as number of BDD nodes) in conjunctive partitioning. The default value is 1000 BDD nodes.


#### Abstract

affinity Environment Variable


This variable controls whether to enables the affinity clustering heuristic as described in [MHS00]. Possible values are 0 or 1 : the default value is 1 .

## trans_order file

## Environment Variable

Reads the a variables list from file $t v$ file, to be used when clustering the transition relation. This feature has been provided by Wendy Johnston, University of Queensland. The results of Johnston's research have been presented at FM 2006 in Hamilton, Canada. See [WJKWLvdBR06].
image_cluster_size Environment Variable

One of the parameters to configure the behaviour of the Iwls95CP partitioning algorithm. image_cluster_size is used as threshold value for the clusters. The default value is 1000 BDD nodes.

## image_W $\{\mathbf{1 , 2 , 3 , 4}\}$

Environment Variable
The other parameters for the Iwls95CP partitioning algorithm. These attribute different weights to the different factors in the algorithm. The default values are $6,1,1,6$ respectively. (For a detailed description, please refer to $\left[\mathrm{RAP}^{+} 95\right]$.)

## iwls95preorder

Environment Variable
Enables cluster preordering following heuristic described in [RAP $\left.{ }^{+} 95\right]$, possible values are 0 or 1 . The default value is 0 . Preordering can be very slow.
image_verbosity Environment Variable

Sets the verbosity for the image method $I w l s 95 C P$, possible values are 0 or 1 . The default value is 0 .

```
print_iwls95options [-h]
```

This command prints out the configuration parameters of the IWLS95 clustering algorithm, i.e. image_verbosity, image_cluster_size and image_W $\{1,2,3,4\}$.
go [-h] [-f]
This command initializes the system for verification. It is equivalent to the command sequence read_model, flatten_hierarchy, encode_variables, build_flat_model, build_model.
If some commands have already been executed, then only the remaining ones will be invoked.


Command Options:
Forces model construction even when Cone Of Influence is enabled.
get_internal_status - Prints out the internal status of the system.

```
get_internal_status [-h]
```

Prints out the internal status of the system. i.e.

- -1: read_model has not yet been executed or an error occurred during its execution.
- 0: flatten_hierarchy has not yet been executed or an error occurred during its execution.
- 1: encode_variables has not yet been executed or an error occurred during its execution.
- 2: build_model has not yet been executed or an error occurred during its execution.
process_model - Performs the batch steps and then returns control to the inter-

```
process_model [-h] [-f] [-r] [-i model-file] [-m Method]
```

Reads the model, compiles it into BDD and performs the model checking of all the specification contained in it. If the environment variable forward_search has been set before, then the set of reachable states is computed. If the option $-r$ is specified, the reordering of variables is performed and a dump of the variable ordering is performed accordingly. This command simulates the batch behavior of NUXMV and then returns the control to the interactive shell.

Command Options:

$$
\begin{array}{ll}
-\mathrm{f} & \text { Forces the model construction even when Cone Of Influence is enabled. } \\
-\mathrm{r} & \begin{array}{l}
\text { Forces a variable reordering at the end of the computation, and dumps the } \\
\text { new variables ordering to the default ordering file. This options acts like the } \\
\text { command line option -reorder. }
\end{array} \\
-\mathrm{i} \text { model-file } & \begin{array}{l}
\text { Sets the environment variable input_file to file model-file, and } \\
\text { reads the model from file model-file. }
\end{array} \\
-\mathrm{m} \text { Method } & \begin{array}{l}
\text { Sets the environment variable partition_method to Method and uses it } \\
\text { as partitioning method. }
\end{array}
\end{array}
$$

```
build_flat_model [-h]
```

Compiles the flattened hierarchy into SEXP (initial states, invariants, and transition relation).

```
build_boolean_model - Compiles the flattened hierarchy into boolean Scalar
```

build_boolean_model [-h] [-f]

```

Compiles the flattened hierarchy into boolean SEXP (initial states, invariants, and transition relation).

Command Options:
\(-£ \quad\) Forces the boolean model construction.
write_flat_model - Writes a flat model to a file
Command
write_flat_model [-h] [-A] [-o filename]
Writes the currently loaded SMV model in the specified file, after having flattened it. Processes are eliminated and a corresponding equivalent model is printed out.

If no file is specified, the file specified via the environment variable output_flatten_model_file is used if any, otherwise standard output is used.

Command Options:
```

-o filename Attempts to write the flat SMV model in filename
-A Writes the flat SMV model using a renaming map to "anonimize" the model.
All the symbols except numerical constanst will be renamed.

```

\section*{output flatten_model file}

The file where the flattened model has to be written. The default value is 'st dout'.

\section*{daggifier_enabled}

Environment Variable
Determines whether the expression daggifier in the model dumping features is enabled or not. The default is enabled.

Environment Variable
Sets the minimum threshold for expressions depth to be daggified.
daggifier_counter_threshold Environment Variable

Sets the minimum threshold for expressions count to be daggified. (i.e. expression must show at least Number time to be daggified

\section*{daggifier_statistics}

Prints daggifier statistics after model dumping.
```

write_boolean_model [-h] [-o filename]

```

Writes the currently loaded NUXMV model in the specified file, after having flattened and booleanized it. Processes are eliminated and a corresponding equivalent model is printed out.

If no file is specified, the file specified via the environment variable output_boolean_model_file is used if any, otherwise standard output is used.

Command Options:
-o filename Attempts to write the flat and boolean NUXMV model in filename

In NUXMV scalar variables are dumped as DEFINEs whose body is their boolean encoding.
This allows the user to still express and see parts of the generated boolean model in terms of the original model's scalar variables names and values, and still keeping the generated model purely boolean.

Also, symbolic constants are dumped within a CONSTANTS statement to declare the values of the original scalar variables' for future reading of the generated file.
When NUXMV detects that there were triggered one or more dynamic reorderings in the BDD engine, the command write_boolean_model also dumps the current variables ordering, if the option output_order_file is set.
The dumped variables ordering will contain single bits or scalar variables depending on the current value of the option write_order_dumps_bits. See command write_order for further information about variables ordering.

\section*{output_boolean_model_file}

Environment Variable
The file where the flattened and booleanized model has to be written. The default value is 'st dout'.

\section*{output_word_format}

Environment Variable
This variable sets in which base unsigned word[•] and signed word \([\bullet]\) constants are outputted (during traces, counterexamples, etc, printing). Possible values are \(2,8,10\) and 16 . Note that if a part of an input file is outputted (for example, if a specification expression is outputted) then the unsigned word \([\bullet]\) and signed word[•] constants remain in same format as they were written in the input file.

\subsection*{4.2 Commands for Checking Specifications}

The following commands allow for the BDD-based model checking of a NUXMV model. These commands can be used only for NUXMV models that do not contain Real or Integers.
compute_reachable - Computes the set of reachable states
```

compute_reachable [-h] [-k number] [-t seconds]

```

Computes the set of reachable states. The result is then used to simplify image and preimage computations. This can result in improved performances for models with sparse state spaces. Sometimes the execution of this command can take much time because the computation of reachable states may be very expensive. Use the -k option to limit the number of forward step to perform. If the reachable states has been already computed the command returns immediately since there is nothing more to compute.

\section*{Command Options:}
-k number If specified, limits the computation of reachable states to perform number steps forward starting from the last computed frontier. This means that you can expand the computed reachable states incrementally using this option.
-t seconds If specified, forces the computation of reachable states to end after "seconds" seconds. This limit could not be precise since the if the computation of a step is running when the limit occurs, the computation is not interrupted until the end of the step

```

print_reachable_states [-h] [-v] [-d] [-f] [-o filename]

```

Prints the number of reachable states of the given model. In verbose mode, prints also the list of all reachable states, if they are less than \(2^{16}\). The reachable states are computed if needed.

Command Options:
\begin{tabular}{ll}
-v & Prints the list of reachable states \\
-d & Prints the list of reachable states with defines (Requires -v ) \\
-f & Prints the formula representing the reachable states \\
-o filename & Prints the result on the specified filename instead of on standard output
\end{tabular}
check fsm - Checks the transition relation for totality.
Command
```

check_fsm [-h] [-m | -o output-file]

```

Checks if the transition relation is total. If the transition relation is not total then a potential deadlock state is shown.

Command Options:
\[
\begin{array}{ll}
-m & \begin{array}{l}
\text { Pipes the output generated by the command to the program specified by the } \\
\text { PAGER shell variable if defined, else through the UNIX command "more". }
\end{array} \\
-\circ \text { output-file } & \text { Writes the output generated by the command to the file output-file. }
\end{array}
\]

At the beginning reachable states are computed in order to guarantee that deadlock states are actually reachable.
```

check fsm Environment Variable

```

Controls the activation of the totality check of the transition relation during the process_model call. Possible values are 0 or 1. Default value is 0 .
```

print_fsm_stats [-h] | [-m] | [-p] | [-o output-file]

```

This command prints out information regarding the fsm and each cluster. In particular for each cluster it prints out the cluster number, the size of the cluster (in BDD nodes), the variables occurring in it, the size of the cube that has to be quantified out relative to the cluster and the variables to be quantified out.
Also the command can print all the normalized predicates the FMS consists of. A normalized predicate is a boolean expression which does not have other boolean sub-expressions. For example, expression ( \(b<0\) ? \(\mathrm{a} / \mathrm{b}: 0)=\mathrm{c}\) is normalized into ( \(\mathrm{b}<0\) ? \(\mathrm{a} / \mathrm{b}=\mathrm{c}: 0=\mathrm{c}\) ) which has 3 normalized predicates inside: \(\mathrm{b}<0, \mathrm{a} / \mathrm{b}=\mathrm{c}, 0=\mathrm{c}\).

Command Options:
\begin{tabular}{ll}
-h & \begin{tabular}{l} 
Prints the command usage. \\
Pipes the output generated by the command to the program specified by the
\end{tabular} \\
-m & \begin{tabular}{l} 
PAGER shell variable if defined, else through the UNIX command "more".
\end{tabular} \\
-p & \begin{tabular}{l} 
Prints out the normalized predicates the FSM consists of. Expressions in \\
properties are ignored.
\end{tabular} \\
-o output-file & \begin{tabular}{l} 
Writes the output generated by the command to the file output-file.
\end{tabular}
\end{tabular}
print_fair_states - Prints out the number of fair states
Command
```

print_fair_states [-h] [-v]

```

Prints the number of fair states of the given model. In verbose mode, prints also the list of all fair states, if they are less than \(2^{16}\).
print fair_transitions - Prints out the number of fair states
Command
```

print_fair_transitions [-h] [-v]

```

Prints the number of fair transitions of the given model. In verbose mode, prints also the list of all fair transitions, if they are less than \(2^{16}\). The transitions are displayed as state-input pairs.

\section*{check_ctlspec - Performs fair CTL model checking.}
```

check_ctlspec [-h] [-m | -o output-file] [-n number | -p
"ctl-expr [IN context]" | -P "name"]

```

Performs fair CTL model checking.
A ctl-expr to be checked can be specified at command line using option -p . Alternatively, option -n can be used for checking a particular formula in the property database. If neither -n nor -p nor -P are used, all the SPEC formulas in the database are checked.
See variable use_coi_size_sorting for changing properties verification order.
Command Options:
-m
-o output-file
-p "ctl-expr [INA CTL formula to be checked. context is the module instance name which context]" the variables in ctl-expr must be evaluated in.
-n number \(\quad\) Checks the CTL property with index number in the property database.
-P name Checks the CTL property named name in the property database.

If the ag_only_search environment variable has been set, then a specialized algorithm to check AG formulas is used instead of the standard model checking algorithms.

Enables the use of an ad hoc algorithm for checking AG formulas. Given a formula of the form AG alpha, the algorithm computes the set of states satisfying alpha, and checks whether it contains the set of reachable states. If this is not the case, the formula is proved to be false.

Enables the computation of the reachable states during the process model command and when used in conjunction with the ag_only_search environment variable enables the use of an ad hoc algorithm to verify invariants. This option is set to true by default.

\section*{Itl_tableau forward_search}

Environment Variable
Forces the computation of the set of reachable states for the tableau resulting from BDD-based LTL model checking, performed by command check_ltlspec. If the variable ltl_tableau_forward_search is not set (default), the resulting tableau will inherit the computation of the reachable states from the model, if enabled (see environment variable use_reachable_states). If the variable is set to true, the set of reachable states will be calculated for the model and for the tableau resulting from LTL model checking.
Remark. This might improve performances of the command check_ltlspec, but may also lead to a dramatic slow down for some modeld. This variable has effect only when the calculation of reachable states for the model is enabled (see forward_search).

The algorithm used to determine language emptiness of a Büchi fair transition system. The algorithm may be used from the following commands: check_ltlspec, check_pslspec. Possible values are:
- EL_bwd The default value. The Emerson-Lei algorithm [EL86] in its usual backwards direction, i.e., using backward image computations.
- ELfwd A variant of the Emerson-Lei algorithm that uses only forward image computations (see, e.g., [HKQ03]). This variant requires the variables forward_search, ltl_tableau_forward_search, use_reachable_states to be set. Furthermore, counterexample computation is not yet implemented, i.e., counter_examples should not be set. When invoking one of the commands mentioned above, all required settings are performed automatically if not already found as needed, and are restored after execution of the command.
```

check_invar [-h] [-m | -o output-file] [-n number | -p
"invar-expr [IN context]" | -P "name"] [-s strategy] [-e
f-b-heuristic] [-j b-b-heuristic] [-t threshold] [-k length]

```

Performs invariant checking on the given model. An invariant is a set of states. Checking the invariant is the process of determining that all states reachable from the initial states lie in the invariant. Invariants to be verified can be provided as simple formulas (without any temporal operators) in the input file via the INVARSPEC keyword or directly at command line, using the option -p .
Option -n can be used for checking a particular invariant of the model. If neither -n nor -p are used, all the invariants are checked.
During checking of invariants all the fairness conditions associated with the model are ignored.
If an invariant does not hold, a proof of failure is demonstrated. This consists of a path starting from an initial state to a state lying outside the invariant. This path has the property that it is the shortest path leading to a state outside the invariant.

A search strategy can be specified with -s option. This is useful to speed up the check in some situations. If "forward-backward" or "bdd-bmc" strategy is specified then it is possible to choose a search heuristic with -e option; "bdd-bmc" strategy has some other options explained below.

See variable use_coi_size_sorting for changing properties verification order.

\section*{Command Options:}
-m
-o output-file
-n number
-p "invar-expr
[IN cont The command line specified invariant formula to be verified. context is
" the module instance name which the variables in invar-expr must be evaluated in.
\(-P\) name Checks the INVAR property named name in the property database.
-s strategy
Pipes the output generated by the program in processing INVARSPEC s to the program specified by the PAGER shell variable if defined, else through the UNIX command "more".

Writes the output generated by the command in processing INVARSPEC s to the file output-file.

Chooses the strategy to use while performing reachability analysis. Possible strategies are:
- "forward" Explore the search space from initial states and try to reach bad states.
- "backward" Explore the search space from bad states and try to reach initial states.
- "forward-backward" Explore the search space using a heuristic to decide at each step whether to move from bad states or from reachable states.
- "bdd-bmc" Explore the search space using BDD with "forwardbackward" strategy and use a heuristic (specified with \(-j\) option) to decide if to switch from BDD technology to BMC. The idea is to expand the sets of states reachable from both bad and initial states, eventually stop and search for a path between frontiers using BMC technology. Options \(-j,-t\) and \(-k\) are enabled only when using this strategy. Note that the algorithm used for the BMC approach is the one specified in the variable "bmc_invar_alg".
If this option is not specified, the value of the environment variable "check_invar_strategy" is considered.
-e f-b-heuristic Specify the heuristic that decides at each step if we must expand reachable states or bad states. This option is enabled only when using "forwardbackward" or "bdd-bmc" strategies. Possible values are "zigzag" and "smallest". "zigzag" forces to perform a step forward and the next step backward and so on, while "smallest" performs a step from the frontier with the BDD representing the state is smaller. If this option is not specified, the value of the environment variable "check_invar_forward_backward_heuristic" is considered.
-j b-b-heuristic When using "bdd-bmc" strategy specify the heuristic that decides at which step we must switch from BDD to BMC technolgy. You should use the option \(-t\) to specify the threshold for the chosen heuristic. Possible heuristics are "steps" and "size". "steps" forces to switch after a number of steps equal to the threshold, while "size" switch when BDDs are bigger (in the number of nodes) than the threshold. If this option is not specified, the value of the environment variable "check_invar_bddbmc_heuristic" is considered.
-t threshold When using "bdd-bmc" strategy specify the threshold for the chosen heuristic. If this option is not specified, the value of the environment variable "check_invar_bddbmc_threshold" is considered.
-k length When using "bdd-bmc" strategy specify the maximum length of the path to search for during BMC search. If this option is not specified, the value of the environment variable "bmc_length" is considered.

Determines default search strategy to be used when using command "check_invar". See the documentation of "check_invar" for a detailed description of possible values and intended semantics.

Determines default forward-backward heuristic to be used when using command "check_invar". See the documentation of "check_invar" for a detailed description of possible values and intended semantics.
check_invar_bdd_bmc_heuristic Environment Variable

Determines default bdd-bmc heuristic to be used when using command "check_invar". See the documentation of "check_invar" for a detailed description of possible values and intended semantics.

\section*{check_invar_bdd_bmc_threshold}

Determines default bdd-bmc threshold to be used when using command "check_invar". See the documentation of "check_invar" for a detailed description of possible values and intended semantics.
check Itlspec - Performs LTL model checking
Command
```

check_ltlspec [-h] [-m | -o output-file] [-n number | -p "ltl-expr [IN
context]" | -P "name" ]

```

Performs model checking of LTL formulas. LTL model checking is reduced to CTL model checking as described in the paper by [CGH97a].
A ltl-expr to be checked can be specified at command line using option -p . Alternatively, option -n can be used for checking a particular formula in the property database. If neither -n nor -p are used, all the LTLSPEC formulas in the database are checked.
See variable use_coi_size_sorting for changing properties verification order.


\section*{Command Options:}
```

-m Pipes the output generated by the command in processing LTLSPECS to the
program specified by the PAGER shell variable if defined, else through the
UNIX command "more".
-o output-file Writes the output generated by the command in processing LTLSPECs to the
file output-file.
-p "ltl-expr AnLTL formula to be checked. context is the module instance name which
[IN context]" the variables in ltl-expr must be evaluated in.
-P "name" Checks the LTL property named name
-n number Checks the LTL property with index number in the property database.

```

Informs the lt12smv tableau constructor to generate a symbolic fair transition system for the given LTL formula with one single Justice constraint instead of possibly more than one. (This is achieved by replacing the multiple Justice with a single Justice plus a an additional monitor.) By default multiple Justice are built.

\section*{check_compute - Performs computation of quantitative characteristics}

Command
```

check_compute [-h] [-m | -o output-file] [-n number | -p
"compute-expr [IN context]" | -P "name"]

```

This command deals with the computation of quantitative characteristics of real time systems. It is able to compute the length of the shortest (longest) path from two given set of states.
```

MAX [ alpha , beta ]
MIN [ alpha , beta ]

```

Properties of the above form can be specified in the input file via the keyword COMPUTE or directly at command line, using option -p .
If there exists an infinite path beginning in a state in start that never reaches a state in final, then infinity is returned. If any of the initial or final states is empty, then undefined is returned.
Option -n can be used for computing a particular expression in the model. If neither -n nor -p are used, all the COMPUTE specifications are computed.

It is important to remark here that if the FSM is not total (i.e. it contains deadlock states) COMPUTE may produce wrong results. It is possible to check the FSM against deadlock states by calling the command check_fsm.

See variable use_coi_size_sorting for changing properties verification order.
Command Options:
\(-m\)
Pipes the output generated by the command in processing COMPUTEs to the program specified by the PAGER shell variable if defined, else through the UNIX command "more".
-o output-file Writes the output generated by the command in processing computes to the file output-file.
-p "compute-expr A CoMPUTE formula to be checked. context is the module instance name [IN context]" which the variables in compute-expr must be evaluated in.
\[
\begin{array}{ll}
-\mathrm{n} \text { number } & \text { Computes only the property with index number. } \\
-\mathrm{P} \text { name } & \text { Checks the COMPUTE property named name in the property database. }
\end{array}
\]
```

check_property - Checks a property into the current list of properties, or a

```
check_property [-h] [-n number | -P "name"] | [(-c | -l | -i | -s | -q )
[-p "formula [IN context]"]]
```

Checks the specified property taken from the property list, or adds the new specified property and checks it. It is possible to check LTL, CTL, INVAR, PSL and quantitative (COMPUTE) properties. Every newly inserted property is inserted and checked.
See variable use_coi_size_sorting for changing properties verification order.

## Command Options:

| -n number | Checks the property stored at the given index |
| :---: | :---: |
| -P name | Checks the property named name in the property database. |
| -c | Checks all the CTL properties not already checked. If -p is used, the given formula is expected to be a CTL formula. |
| -1 | Checks all the LTL properties not already checked. If -p is used, the given formula is expected to be a LTL formula. |
| -i | Checks all the INVAR properties not already checked. If -p is used, the given formula is expected to be a INVAR formula. |
| -s | Checks all the PSL properties not already checked. If -p is used, the given formula is expected to be a PSL formula. |
| -q | Checks all the COMPUTE properties not already checked. If -p is used, the given formula is expected to be a COMPUTE formula. |
| -p "formula [IN con | Checks the formula specified on the command-line. context is the module instance name which the variables in formula must be evaluated in. |

```
add_property [-h] [(-c | -l | -i | -q | -s) -p "formula
[IN context]"] [-n "name"]
```

Adds a property in the list of properties. It is possible to insert LTL, CTL, INVAR, PSL and quantitative (COMPUTE) properties. Every newly inserted property is initialized to unchecked. A type option must be given to properly execute the command.

Command Options:

| $-c$ | Adds a CTL property. |
| :--- | :--- |
| -1 | Adds an LTL property. |
| $-i$ | Adds an INVAR property. |
| $-s$ | Adds a PSL property. |
| $-q$ | Adds a quantitative (COMPUTE) property. |

-p "formula Adds the formula specified on the command-line. context is the module [IN context]" instance name which the variables in formula must be evaluated in.
-n "name" Sets the name of the property to "name"

```
show_property [-h] [-n idx | -P "name"] [-c | -l | -i | -s | -q] [-f |
-v | -u] [-m | -o] [-F format]
```

Shows the properties currently stored in the list of properties. This list is initialized with the properties (CTL, LTL, INVAR, COMPUTE) present in the input file, if any; then all of the properties added by the user with the relative check_property or add_property commands are appended to this list. For every property, the following informations are displayed:

- the identifier of the property (a progressive number);
- the property name if available;
- the property formula;
- the type (CTL, LTL, INVAR, PSL, COMPUTE)
- the status of the formula (Unchecked, True, False) or the result of the quantitative expression, if any (it can be infinite);
- if the formula has been found to be false, the index number of the corresponding counterexample trace.

By default, all the properties currently stored in the list of properties are shown. Specifying the suitable options, properties with a certain status (Unchecked, True, False) and/or of a certain type (e.g. CTL, LTL), or with a given identifier, it is possible to let the system show a restricted set of properties. It is allowed to insert only one option per status and one option per type.

## Command Options:

| -P name | Prints out the property named "name" |
| :---: | :---: |
| -n idx | Prints out the property numbered "idx" |
| -C | Prints only CTL properties |
| -1 | Prints only LTL properties |
| -i | Prints only INVAR properties |
| -q | Prints only COMPUTE properties |
| -u | Prints only unchecked properties |
| -t | Prints only those properties found to be true |
| -f | Prints only those properties found to be false |
| -s | Prints the number of stored properties |
| -o filename | Writes the output generated by the command to filename |
| -F format | Prints with the specified format. tabular and $x m l$ are common formats, however use $-F$ help to see all available formats. |
| -m | Pipes the output through the program specified by the PAGER shell variable if defined, else through the UNIX "more" command |

$\underset{\substack{\text { FoNDOZIONE } \\ \text { BRUNO KESSER }}}{ }$
write_coi_model - Writes a restricted flat model to a file

```
write_coi_model [-h] [-n idx | -p "expr" | -P "name"] [-c | -l | -i | -s
| -q] [-C] [-g]
```

Writes the currently loaded SMV model in the specified file, after having flattened it. If a property is specified, the dumped model is the result of applying the Cone Of Influence over that property. otherwise, a restricted SMV model is dumped for each property in the property database.
Processes are eliminated and a corresponding equivalent model is printed out.
If no file is specified, stderr is used for output
Command Options:

| -0 filename | Attempts to write the flat SMV model in filename |
| :--- | :--- |
| -p expr | Applies COI for the given expression expression. Notice that also the prop- <br> erty type has to be specified |
| -P name | Applies COI for property named "name" |
| -n idx | Applies COI for property stored with index "idx" |
| -c | Dumps COI model for all CTL properties |
| -l | Dumps COI model for all LTL properties |
| -i | Dumps COI model for all INVAR properties |
| -s | Dumps COI model for all PSL properties |
| -q | Dumps COI model for all COMPUTE properties |
| -C | Only prints the list of variables that are in the COI of properties |
| -g | Dumps the COI model that represents the union of all COI properties |

Uses the cone of influence reduction when checking properties. When cone of influence reduction is active, the problem encoded in the solving engine consists only of the relevant parts of the model for the property being checked. This can greatly help in reducing solving time and memory usage. Note however, that a COI counter-example trace may or may not be a valid counter-example trace for the original model.

## use_coi_size_sorting

Environment Variable
Uses the cone of influence variables set size for properties sorting, before the verification step. If set to 1 , properties are verified starting with the one that has the smallest COI set, ending with the property with the biggest COI set. If set to 0 , properties are verified according to the declaration order in the input file

Determines how properties are printed. The following methods are available:
name. Prints the property name. If not available, defaults to method "index".
index. Prints the property index. If not available, defaults to method "truncated".
truncated. Prints the formula of the property. If the formula is longer than 40 characters, it is truncated.
formula. The default method, simply prints the formula.

### 4.3 Commands for Bounded Model Checking

In this section we describe in detail the commands for doing and controlling Bounded Model Checking in NUXMV. Bounded Model Checking is based on the reduction of the bounded model checking problem to a propositional satisfiability problem. After the problem is generated, nUXmV internally calls a propositional SAT solver in order to find an assignment which satisfies the problem. Currently nuXmv supplies two SAT solvers: Zchaff and MiniSat. If none of the two is enabled, all Bounded Model Checking part in nUXmV will not be available. Notice that Zchaff and MiniSat are for non-commercial purposes only. They are therefore not included in the source code distribution or in some of the binary distributions of NUXMV.

Some commands for Bounded Model Checking use incremental algorithms. These algorithms exploit the fact that satisfiability problems generated for a particular bounded model checking problem often share common subparts. So information obtained during solving of one satisfiability problem can be used in solving of another one. The incremental algorithms usually run quicker then non-incremental ones but require a SAT solver with incremental interface. At the moment, only Zchaff and MiniSat offer such an interface. If none of these solvers are linked to NUXmv, then the commands which make use of the incremental algorithms will not be available.

It is also possible to generate the satisfiability problem without calling the SAT solver. Each generated problem is dumped in DIMACS format to a file. DIMACS is the standard format used as input by most SAT solvers, so it is possible to use NUXMV with a separate external SAT solver. At the moment, the DIMACS files can be generated only by commands which do not use incremental algorithms.
bmc_setup - Builds the model in a Boolean Epression format.

```
bmc_setup [-h]
```

You must call this command before use any other bmc-related command. Only one call per session is required.

```
go_bmc [-h] [-f]
```

This command initializes the system for verification. It is equivalent to the command sequence read_model, flatten_hierarchy, encode_variables, build_boolean_model, bmc_setup. If some commands have already been executed, then only the remaining ones will be invoked.

Command Options:

This variable enables the Sexp inlining when the boolean model is built. Sexp inlining is performed in a similar way to RBC inlining (see system variable rbc_inlining) but the underlying structures and kind of problem are different, because inlining is applied at the Sexp level instead of the RBC level.
Inlining is applied to initial states, invariants and transition relations. By default, Sexp inlining is disabled.

When set, this variable makes BMC perform the RBC inlining before committing any problem to the SAT solver. Depending on the problem structure and length, the inlining may either make SAT solving much faster, or slow it down dramatically. Experiments showed an average improvement in time of SAT solving when RBC inlining is enabled. RBC inlining is enabled by default.

The idea about inlining was taken from [ABE00] by Parosh Aziz Abdulla, Per Bjesse and Niklas Eén.

## rbc_rbc2cnf_algorithm

Environment Variable
This variable defines the algorithm used for conversion from RBC to CNF format in which a problem is supplied to a SAT solver. The default value 'sheridan' refers to [She04] algorithm which allows to obtain a more compact CNF formulas. The other value 'tseitin' refers to a standard Tseiting transformation algorithm.
check_ltlspec_bmc - Checks the given LTL specification, or all LTL specifica-
Command tions if no formula is given. Checking parameters are the maximum length and the loopback value

```
check_ltlspec_bmc [-h ] | [-n idx | -p "formula [IN context]" | -P
"name"] [-k max_length] [-l loopback] [-o filename]
```

This command generates one or more problems, and calls SAT solver for each one. Each problem is related to a specific problem bound, which increases from zero ( 0 ) to the given maximum problem length. Here max_length is the bound of the problem that system is going to generate and solve. In this context the maximum problem bound is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length. The single generated problem also depends on the loopback parameter you can explicitly specify by the -l option, or by its default value stored in the environment variable bmc_loop.back.

The property to be checked may be specified using the $-n \quad i d x$ or the -p "formula" options. If you need to generate a DIMACS dump file of all generated problems, you must use the option -o "filename".

## Command Options:

```
-n index index is the numeric index of a valid LTL specification formula actually lo-
    cated in the properties database.
-p "formula Checks the formula specified on the command-line. context is the mod-
    [IN context]" ule instance name which the variables in formula must be evaluated in.
-P name Checks the LTL property named name in the property database.
-k max_length max_length is the maximum problem bound to be checked. Only natural
    numbers are valid values for this option. If no value is given the environment
    variable bmc_length is considered instead.
-1 loopback
-o filename
The loopback value may be:
- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- a negative number in ( -1 , -bmc_length). In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' \(x\) ', which means "no loopback".
- the symbol ' \(\star\) ', which means "all possible loopbacks from zero to lengthl".
-o filename filename is the name of the dumped dimacs file. It may contain special symbols which will be macro-expanded to form the real file name. Possible symbols are:
```

- @F: model name with path part.
- @f: model name without path part.
- @k: current problem bound.
- @1: current loopback value.
- @ n : index of the currently processed formula in the property database.
- @ @: the ‘@' character.
check_Itlspec_bmc_onepb - Checks the given LTL specification, or all LTL
specifications if no formula is given. Checking parameters are the single prob-
lem bound and the loopback value

```
check_ltlspec_bmc_onepb [-h ] | [ -n idx | -p "formula" [IN context] | -P
"name"] [-k length] [-l loopback] [-o filename]
```

As command check_ltlspec_bmc but it produces only one single problem with fixed bound and loopback values, with no iteration of the problem bound from zero to max_length.

## Command Options:



- @F: model name with path part.
- @f: model name without path part.
- @k: current problem bound.
- @1: current loopback value.
- @ n : index of the currently processed formula in the property database.
- @ @: the '@' character.

[^8]

```
gen_ltlspec_bmc [-h] | [ -n idx | -p "formula" [IN context] | -P "name"]
[-k max_length] [-l loopback] [-o filename]
```

This command generates one or more problems, and dumps each problem into a dimacs file. Each problem is related to a specific problem bound, which increases from zero (0) to the given maximum problem bound. In this short description length is the bound of the problem that system is going to dump out.
In this context the maximum problem bound is represented by the max_length parameter, or by its default value stored in the environment variable bmc_length.
Each dumped problem also depends on the loopback you can explicitly specify by the -1 option, or by its default value stored in the environment variable bmc_loopback.
The property to be checked may be specified using the -n idx or the -p "formula " options.
You may specify dimacs file name by using the option -o filename, otherwise the default value stored in the environment variable bmc_dimacs_filename will be considered.

Command Options:
> -n index
> -p "formula Checks the formula specified on the command-line. context is the mod-
> [IN context]" ule instance name which the variables in formula must be evaluated in.
> -P name $\quad$ Checks the LTL property named name in the property database.
> -k max_length max_length is the maximum problem bound used when increasing problem bound starting from zero. Only natural numbers are valid values for this
option. If no value is given the environment variable bmc_length value is bound starting from zero. Only natural numbers are valid values for this
option. If no value is given the environment variable bmc_length value is considered instead.
> -1 loopback The loopback value may be:
> index is the numeric index of a valid LTL specification formula actually located in the properties database. The validity of index value is checked out by the system.

- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of bound and loopback will be skipped during the generation and dumping process.
- a negative number in ( $-1,-b m c$ _length $)$. In this case loopback is considered a value relative to max_length. Any invalid combination of bound and loopback will be skipped during the generation process.
- the symbol ' $X$ ', which means "no loopback".
- the symbol ' $\star$ ', which means 'all possible loopback from zero to length-l".
-o filename
filename is the name of dumped dimacs files. If this options is not specified, variable bmc_dimacs_filename will be considered. The file name string may contain special symbols which will be macro-expanded to form the real file name. Possible symbols are:
- @F: model name with path part.
- @f: model name without path part.
- @k: current problem bound.
- @1: current loopback value .
- @ n : index of the currently processed formula in the property database.
- @ @: the '@' character.

```
gen_ltlspec_bmc_onepb [-h ] | [ -n idx | -p "formula" [IN context] | -P
"name"] [-k length] [-l loopback] [-o filename]
```

As the gen_ltlspec_bmc command, but it generates and dumps only one problem given its bound and loopback.

Command Options:

| $-n$ index | index is the numeric index of a valid LTL specification formula actually lo- <br> cated in the properties database. The validity of index value is checked out <br> by the system. |
| :--- | :--- |
| -p "formula | Checks the formula specified on the command-line. context is the mod- <br> [IN context] <br> ule instance name which the variables in formula must be evaluated in. <br> -P name |
| -l length | Checks the LTL property named name in the property database. <br> length is the single problem bound used to generate and dump it. Only natu- <br> ral numbers are valid values for this option. If no value is given the environ- <br> ment variable bmc_length is considered instead. |
| -1 loopback | The loopback value may be: |

- a natural number in ( 0 , length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation and dumping process.
- negative number in ( -1 , -length). Any invalid combination of length and loopback will be skipped during the generation process.
- the symbol ' $X$ ', which means "no loopback".
- the symbol ' $\star$ ', which means "all possible loopback from zero to length-l".
-o filename filename is the name of the dumped dimacs file. If this options is not specified, variable bmc_dimacs_filename will be considered. The file name string may contain special symbols which will be macro-expanded to form the real file name. Possible symbols are:
- @F: model name with path part
- @f: model name without path part
- @k: current problem bound
- @1: current loopback value
- @ n : index of the currently processed formula in the property database
- @ @: the '@' character
checkltlspec_bmc_inc - Checks the given LTL specification, or all LTL spec-

```
check_ltlspec_bmc_inc [-h ] | [-n idx | -p "formula [IN context]" | -P
"name" ] [-k max_length] [-l loopback]
```

For each problem this command incrementally generates many satisfiability subproblems and calls the SAT solver on each one of them. The incremental algorithm exploits the fact that subproblems have common subparts, so information obtained during a previous call to the SAT solver can be used in the consecutive ones. Logically, this command does the same thing as check_ltlspec_bmc (see the description on page 68) but usually runs considerably quicker. A SAT solver with an incremental interface is required by this command, therefore if no such SAT solver is provided then this command will be unavailable.
See variable use_coi_size_sorting for changing properties verification order.

## Command Options:

```
-n index
```

index is the numeric index of a valid LTL specification formula actually located in the properties database.
-p "formula Checks the formula specified on the command-line. context is the mod-
[IN context]" ule instance name which the variables in formula must be evaluated in.
-P name Checks the LTL property named name in the property database.

- k max_length max_length is the maximum problem bound must be reached. Only natural numbers are valid values for this option. If no value is given the environment variable bmc_length is considered instead.
-l loopback The loopback value may be:
- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- a negative number in ( $-1,-$ bmc_length). In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' $x$ ', which means "no loopback".
- the symbol ' $\star$ ', which means "all possible loopback from zero to length-l"
mc - Checks the

```
check_ltlspec_sbmc [-h] | [-n idx | -p "formula [IN context]" | -P
"name"] [-k max_length] [-l loopback] [-o filename]
```

This command generates one or more problems, and calls SAT solver for each one. The BMC encoding used is the one by of Latvala, Biere, Heljanko and Junttila as described in [LBHJ05]. Each problem is related to a specific problem bound, which increases from zero ( 0 ) to the given maximum problem length. Here max_length is the bound of the problem that system is going to generate and solve. In this context the maximum problem bound is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length. The single generated problem also depends on the loopback parameter you can explicitly specify by the -1 option, or by its default value stored in the environment variable bmc_loop.back.
The property to be checked may be specified using the $-n i d x$ or the -p "formula" options. If you need to generate a DIMACS dump file of all generated problems, you must use the option -o "filename".

See variable use_coi_size_sorting for changing properties verification order.

## Command Options:

| -n index | index is the numeric index of a valid LTL specification formula actually located in the properties database. |
| :---: | :---: |
| "formula [IN con | Checks the formula specified on the command-line. context is the mod" ule instance name which the variables in formula must be evaluated in. |
| -P name | Checks the LTL property named name in the property database. |
| - k max_length | max_length is the maximum problem bound to be checked. Only natural numbers are valid values for this option. If no value is given the environment variable bmc_length is considered instead. |
| -1 loopback | The loopback value may be: |

- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- a negative number in ( $-1,-b m c$ _length $)$. In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' $x$ ', which means "no loopback".
- the symbol ' $*$ ', which means "all possible loopbacks from zero to lengthl".
-○ filename
filename is the name of the dumped dimacs file. It may contain special symbols which will be macro-expanded to form the real file name. Possible symbols are:
- @F: model name with path part.
- @f: model name without path part.
- @k: current problem bound.
- @1: current loopback value.
- @ n : index of the currently processed formula in the property database.
- @ @: the ‘@' character.

> check_ltlspec_sbmc_inc - Checks the given LTL specification, or all LTL specifications if no formula is given. Checking parameters are the maximum length and the loopack value

```
check_ltlspec_sbmc_inc [-h ] | [ -n idx | -p "formula [IN context]" | -P
"name"] [-k max_length] [-N] [-c]
```

This command generates one or more problems, and calls SAT solver for each one. The Incremental BMC encoding used is the one by of Heljanko, Junttila and Latvala, as described in [KHL05]. For each problem this command incrementally generates many satisfiability subproblems and calls the SAT solver on each one of them. Each problem is related to a specific problem bound, which increases from zero (0) to the given maximum problem length. Here max_length is the bound of the problem that system is going to generate and solve. In this context the maximum problem bound is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length.

The property to be checked may be specified using the $-n$ idx, the -p "formula" or the -P "name" options.

See variable use_coi_size_sorting for changing properties verification order.
Command Options:


```
gen_ltlspec_sbmc [-h ] | [ -n idx | -p "formula [IN context]" | -P
"name" ] [-k max_length] [-l loopback] [-o filename]
```

This command generates one or more problems, and dumps each problem into a dimacs file. The BMC encoding used is the one by of Latvala, Biere, Heljanko and Junttila as described in [LBHJ05]. Each problem is related to a specific problem bound, which increases from zero (0) to the given maximum problem length. Here max_length is the bound of the problem that system is going to generate and dump. In this context the maximum problem bound is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length. The single generated problem also depends on the loopback parameter you can explicitly specify by the -1 option, or by its default value stored in the environment variable bmc_loopback.
The property to be used for the problem dumping may be specified using the -n idx or the -p "formula" options. You may specify dimacs file name by using the option -o "filename", otherwise the default value stored in the environment variable bmc_dimacs_filename will be considered.

## Command Options:

| -n index | index is the numeric index of a valid LTL specification formula actually lo- <br> cated in the properties database. |
| :--- | :--- |
| -p "formula | Dumps the formula specified on the command-line. context is the mod- <br> [IN context] <br> -p <br> " name instance name which the variables in formula must be evaluated in. |
| -k max_length | Checks the LTL property named name <br> max_length is the maximum problem bound to be generated. Only natural <br> numbers are valid values for this option. If no value is given the environment |
| variable bmc_length is considered instead. |  |

- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- a negative number in ( -1, -bmc_length). In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' $x$ ', which means "no loopback".
- the symbol ' $\star$ ', which means "all possible loopbacks from zero to lengthl".
-○ filename filename is the name of the dumped dimacs file. It may contain special symbols which will be macro-expanded to form the real file name. Possible symbols are:
- @F: model name with path part.
- @f: model name without path part.
- @k: current problem bound.
- @1: current loopback value.
- @ n : index of the currently processed formula in the property database.
- @ @: the ‘@' character.

Sets the generated problem bound. Possible values are any natural number, but must be compatible with the current value held by the variable bmc_loopback. The default value is 10 .

Sets the generated problem loop. Possible values are:

- Any natural number, but less than the current value of the variable bmc_length. In this case the loop point is absolute.
- Any negative number, but greater than or equal to -bmc_length. In this case specified loop is the loop length.
- The symbol ' $X$ ', which means "no loopback".
- The symbol ' $\star$ ', which means "any possible loopbacks".

The default value is *.

Uses depth1 optimization for LTL Tableau construction in BMC.
bmc_force_pltl_tableau Environment Variable

Forces to use PLTL instead of LTL for BMC tableau construction.

## bmc_dimacs_filename <br> Environment Variable

This is the default file name used when generating DIMACS problem dumps. This variable may be taken into account by all commands which belong to the gen_ltlspec_bmc family. DIMACS file name can contain special symbols which will be expanded to represent the actual file name. Possible symbols are:

- @F The currently loaded model name with full path.
- @f The currently loaded model name without path part.
- @n The numerical index of the currently processed formula in the property database.
- @k The currently generated problem length.
- @I The currently generated problem loopback value.
- @ @ The '@’ character.

The default value is "@f_k@k_l@l_n@n".

## bmc_sbmc_gf_fg_opt <br> Environment Variable

Controls whether the system exploits an optimization when performing SBMC on formulae in the form $F G p$ or $G F p$. The default value is 1 (active).
check_invar_bmc - Generates and solves the given invariant, or all invariants
Command
if no formula is given

```
check_invar_bmc [-h | -n idx | -p "formula" [IN context] | -P "name"]
[-a alg] [-e] [-k bmc_bound] [-o filename]
```

In Bounded Model Checking, invariants are proved using induction. For this, satisfiability problems for the base and induction step are generated and a SAT solver is invoked on each of them. At the moment, two algorithms can be used to prove invariants. In one algorithm, which we call "classic", the base and induction steps are built on one state and one transition, respectively. Another algorithm, which we call "een-sorensson" [ESO4], can build the base and induction steps on many states and transitions. As a result, the second algorithm is more powerful.
Also, notice that during checking of invariants all the fairness conditions associated with the model are ignored.
See variable use_coi_size_sorting for changing properties verification order.


Command Options:

| -n index | index is the numeric index of a valid INVAR specification formula actually located in the property database. The validity of index value is checked out by the system. |
| :---: | :---: |
| -p "formula <br> [IN con | Checks the formula specified on the command-line. context is the module instance name which the variables in formula must be evaluated in. |
| -P name | Checks the INVAR property named name in the property database. |
| - k max_length | max_length is the maximum problem bound that can be reached. Only natural numbers are valid values for this option. Use this option only if the "een-sorensson" algorithm is selected. If no value is given the environment variable bmc_length is considered instead. |
| -e | Performs an extra induction step for finding a proof. Can be used only with the "een-sorensson" algorithm |
| -a alg | alg specifies the algorithm. The value can be classic or een-sorensson. If no value is given the environment variable bmc_invar_alg is considered instead. |
| -○ filename | filename is the name of the dumped dimacs file. It may contain special symbols which will be macro-expanded to form the real file name. Possible | symbols are:

- @F: model name with path part
- @f: model name without path part
- @n: index of the currently processed formula in the properties database
- @ @: the '@' character
gen_invar_bmc - Generates the given invariant, or all invariants if no formula

```
gen_invar_bmc [-h | -n idx | -p "formula [IN context]" | -P "name"] [-o
filename]
```

At the moment, the invariants are generated using "classic" algorithm only (see the description of check_invar_bmc on page 75).

Command Options:
-n index index is the numeric index of a valid INVAR specification formula actually located in the property database. The validity of index value is checked out by the system.
-p "formula Checks the formula specified on the command-line. context is the mod-
[IN context]" ule instance name which the variables in formula must be evaluated in.
$-P$ name Checks the INVAR property named name in the property database.
-o filename filename is the name of the dumped dimacs file. If you do not use this option the dimacs file name is taken from the environment variable bmc_invar_dimacs_filename.
File name may contain special symbols which will be macro-expanded to form the real dimacs file name. Possible symbols are:

- @F: model name with path part
- @f: model name without path part
- @ $\mathbf{n}$ : index of the currently processed formula in the properties database
- @ @: the '@' character
check_invar_bmc_inc - Generates and solves the given invariant, or all invari- Command
ants if no formula is given, using incremental algorithms
ants if no formula is given, using incremental algorithms

```
check_invar_bmc_inc [-h ] | [ -n idx | -p "formula" [IN context] | -P
"name" ]] [-a algorithm]
```

This command does the same thing as check_invar_bmc (see the description on page 75) but uses an incremental algorithm and therefore usually runs considerably quicker. The incremental algorithms exploit the fact that satisfiability problems generated for a particular invariant have common subparts, so information obtained during solving of one problem can be used in solving another one. A SAT solver with an incremental interface is required by this command. If no such SAT solver is provided then this command will be unavailable.
There are two incremental algorithms which can be used: "Dual" and "ZigZag". Both algorithms are equally powerful, but may show different performance depending on a SAT solver used and an invariant being proved. At the moment, the "Dual" algorithm cannot be used if there are input variables in a given model. For additional information about algorithms, consider [ES04].
Also, notice that during checking of invariants all the fairness conditions associated with the model are ignored.
See variable use_coi_size_sorting for changing properties verification order.

## Command Options:

$$
\begin{array}{ll}
\text {-n index } & \begin{array}{l}
\text { index is the numeric index of a valid INVAR specification formula actually } \\
\text { located in the property database. The validity of index value is checked out } \\
\text { by the system. }
\end{array} \\
\text {-p "formula } & \begin{array}{l}
\text { Checks the formula specified on the command-line. context is the mod- } \\
\text { [IN context] } \\
\text { ule instance name which the variables in formula must be evaluated in. }
\end{array} \\
-\mathrm{P} \text { "name" } & \begin{array}{l}
\text { Checks the INVARSPEC property named name } \\
\text { max_length is the maximum problem bound that can be reached. Only natu- } \\
\text { ral numbers are valid values for this option. If no value is given the environ- } \\
\text { ment variable bmc_length is considered instead. }
\end{array} \\
\text {-a alg } & \begin{array}{l}
\text { alg specifies the algorithm to use. The value can be dual or zigzag. If } \\
\text { no value is given the environment variable } b m c \_i n c \_i n v a r \_a l g ~ i s ~ c o n s i d e r e d ~
\end{array} \\
\text { instead. }
\end{array}
$$

## bmc_invar_alg

Environment Variable
Sets the default algorithm used by the command check_invar_bmc. Possible values are classic and een-sorensson. The default value is classic.

## bmc_inc_invar_alg

Environment Variable
Sets the default algorithm used by the command check_invar_bmc_inc. Possible values are dual and zigzag. The default value is dual.

This is the default file name used when generating DIMACS invar dumps. This variable may be taken into account by the command gen_invar_bmc. DIMACS file name can contain special symbols which will be expanded to represent the actual file name. Possible symbols are:

- @F The currently loaded model name with full path.
- @f The currently loaded model name without path part.
- @ $\mathbf{n}$ The numerical index of the currently processed formula in the properties database.
- @ @ The ‘@’ character.

The default value is "@f_invar_n@n".

## sat_solver

Environment Variable
The SAT solver's name actually to be used. Default SAT solver is MiniSat. Depending on the NuXmy configuration, also the Zchaff SAT solver can be available or not. Notice that Zchaff and MiniSat are for non-commercial purposes only. If no SAT solver has been configured, BMC commands and environment variables will not be available.

```
bmc_pick_state [-h] [-v] [-c "constraint" | -s trace.state] [-r | -i
[-a]]
```

Chooses an element from the set of initial states, and makes it the current state (replacing the old one). The chosen state is stored as the rst state of a new trace ready to be lengthened by steps states by the bmc_simulate command or the bmc_inc_simulate command.

Command Options:

```
-v Verbosely prints the generated trace
_ c constraint Set a constraint to narrow initial states.
-s state Picks state from trace.state label.
-r Randomly picks a state from the set of initial states.
-i Enters simulation's interactive mode.
-a Displays all the state variables (changed and unchanged) in the interactive
session
```

```
omc_simulate [-h] [-p | -v] [-r] [[-c "constraints"] | [-t
"constraints"] ] [-k steps]
```

bmc_simulate does not require a specification to build the problem, because only the model is used to build it. The problem length is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length.

## Command Options:

```
-p Prints the generated trace (only changed variables).
-v Prints the generated trace (all variables).
```

| -r | Pi |
| :---: | :---: |
| - c constraint | Performs a simulation in which computation is restricted to states satisfying those constraints. The desired sequence of states could not exist if such constraints were too strong or it may happen that at some point of the simulation a future state satisfying those constraints doesn't exist: in that case a trace with a number of states less than steps trace is obtained. Note: constraints must be enclosed between double quotes " ". The expression cannot contain next operators, and is automatically shifted by one state in order to constraint only the next steps |
| -t "constraints" | Performs a simulation in which computation is restricted to states satisfying those constraints. The desired sequence of states could not exist if such constraints were too strong or it may happen that at some point of the simulation a future state satisfying those constraints doesn't exist: in that case a trace with a number of states less than steps trace is obtained. Note: constraints must be enclosed between double quotes " ". The expression can contain next operators, and is NOT automatically shifted by one state as done with option -c |
| -k steps | Maximum length of the path according to the constraints. The length of a trace could contain less than steps states: this is the case in which simulation stops in an intermediate step because it may not exist any future state satisfying those constraints. The default value is determined by the default_simulation_steps environment variable |

bmc_inc_simulate - Generates a trace of the model from 0 (zero) to $k$
Command
bmc_inc_simulate [-h] [-p | -v] [-r | -i [-a]] [[-c "constraints"] | [-t "constraints"] ] [-k steps]
Performs incremental simulation of the model. bmc_inc_simulate does not require a specification to build the problem, because only the model is used to build it. The problem length is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length.

Command Options:

| -p | Prints the generated trace (only changed variables). |
| :--- | :--- |
| -v | Prints the generated trace (all variables). |
| -r | Picks a state from a set of possible future states in a random way. |
| -i | Enters simulation's interactive mode. |
| -a | Displays all the state variables (changed and unchanged) in the interactive <br> session |
| -c constraint | Performs a simulation in which computation is restricted to states satisfy- <br> ing those constraints. The desired sequence of states could not exist if |
|  | such constraints were too strong or it may happen that at some point of the <br> simulation a future state satisfying those constraints doesn't exist: in that <br> case a trace with a number of states less than steps trace is obtained. Note: |
|  | constraints must be enclosed between double quotes " ". The expres- <br> sion cannot contain next operators, and is automatically shifted by one state <br> in order to constraint only the next steps |
|  |  |


| $-t$ "constraints" | Performs a simulation in which computation is restricted to states satisfy- <br> ing those constraints. The desired sequence of states could not exist if <br> such constraints were too strong or it may happen that at some point of the <br> simulation a future state satisfying those constraints doesn't exist: in that |
| :--- | :--- |
|  | case a trace with a number of states less than steps trace is obtained. Note: |
| constraints must be enclosed between double quotes " ". The expres- |  |
| sion can contain next operators, and is NOT automatically shifted by one |  |
| state as done with option -c |  |

bmc_simulate_check_feasible_constraints - Checks feasability for the given

Command
constraints
bmc_simulate_check_feasible_constraints [-h] [-q] [-c "constr"]
Checks if the given constraints are feasible for BMC simulation.
Command Options:

```
-q Prints the output in compact form.
-c constr Specify one constraint whose feasability has to be checked (can be used
    multiple times, order is important to read the result)
```


### 4.4 Commands for checking PSL specifications

The following commands allow for model checking of PSL specifications.
check_pslspec - Performs bdd-based PSL model checking

```
check_pslspec [-h] [-m | -o output-file] [-n number | -p
"psl-expr [IN context]" | -P "name"]
```

Check psl properties using bdd-based model checking.
A psl-expr to be checked can be specified at command line using option -p. Alternatively, option $-n$ can be used for checking a particular formula in the property database. If neither -n nor -p are used, all the PSLSPEC formulas in the database are checked.

See variable use_coi_size_sorting for changing properties verification order.
Command Options:
$-m$

- o output-file

Pipes the output generated by the command in processing PSLSPECS to the program specified by the PAGER shell variable if defined, else through the UNIX command "more".
Writes the output generated by the command in processing PSLSPEC s to the file output-file
-p "psl-expr A PSL formula to be checked. context is the module instance name which [IN context]" the variables in psl-expr must be evaluated in.

-n number Checks the PSL property with index number in the property database.<br>$-P$ name Checks the PSL property named name in the property database.

check_pslspec_bmc - Performs SAT-based PSL model checking
Command

```
check_pslspec_bmc [-h] [-m | -o output-file] [-n number | -p
"psl-expr [IN context]" | -P "name"] [-g] [-1] [-k
bmc_lenght] [-l loopback]
```

Check psl properties using SAT-based model checking.
A psl-expr to be checked can be specified at command line using option -p . Alternatively, option -n can be used for checking a particular formula in the property database. If neither -n nor -p are used, all the PSLSPEC formulas in the database are checked. Options -k and -1 can be used to define the maximum problem bound, and the value of the loopback for the single generated problems respectively; their values can be stored in the environment variables bmc_lenght and bmc_loopback. Single problems can be generated by using option -1 . Bounded model checking problems can be generated and dumped in a file by using option -g.
See variable use_coi_size_sorting for changing properties verification order.
Command Options:

| -m | Pipes the output generated by the command in processing PSLSPECS to the program specified by the PAGER shell variable if defined, else through the UNIX command "more". |
| :---: | :---: |
| -○ output-file | Writes the output generated by the command in processing PSLSPEC s to the file output-file |
| $\begin{aligned} & \text {-p "psl-expr } \\ & {[\text { IN context] }} \end{aligned}$ | A PSL formula to be checked. context is the module instance name which the variables in psl-expr must be evaluated in. |
| -n number | Checks the PSL property w |
| -P name | Checks the PSL property named name in the property database. |
| -g | Dumps DIMACS version of bounded model checking problem into a file whose name depends on the system variable bmc_dimacs_filename. This feature is not allowed in combination of the option -i. |
| -1 | Generates a single bounded model checking problem with fixed bound and loopback values, it does not iterate incrementing the value of the problem bound. |
| -k bmc_length | $b m c$ _length is the maximum problem bound to be checked. Only natural numbers are valid values for this option. If no value is given the environment variable bmc_length is considered instead. |
| 1 loopback | The loopback value m |

- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- a negative number in (-1, -bmc_length). In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' $x$ ', which means "no loopback".
- the symbol ' $\star$ ', which means "all possible loopbacks from zero to length$l$ ". If no value is given the environment variable bmc_loopback is considered instead.

```
check_pslspec_bmc_inc [-h] [-m | -o output-file] [-n number | -p
"psl-expr [IN context]" | -P "name"] [-1] [-k
bmc_lenght] [-l loopback]
```

Check psl properties using incremental SAT-based model checking.
A psl-expr to be checked can be specified at command line using option -p . Alternatively, option -n can be used for checking a particular formula in the property database. If neither -n nor -p are used, all the PSLSPEC formulas in the database are checked. Options -k and -1 can be used to define the maximum problem bound, and the value of the loopback for the single generated problems respectively; their values can be stored in the environment variables bmc_lenght and bmc_loopback. Single problems can be generated by using option -1 . Bounded model checking problems can be generated and dumped in a file by using option -g.
See variable use_coi_size_sorting for changing properties verification order.

## Command Options:

-m
-o output-file Writes the output generated by the command in processing PSLSPEC s to the file output-file
-p "psl-expr [IN context]" the variables in psl-expr must be evaluated in.
-n number Checks the PSL property with index number in the property database.
-P name $\quad$ Checks the PSL property named name in the property database.
-1
-k bmc_length bmc_length is the maximum problem bound to be checked. Only natural numbers are valid values for this option. If no value is given the environment variable bmc_length is considered instead.
-1 loopback The loopback value may be:

- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- a negative number in ( $-1,-b m c$ _length). In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' $x$ ', which means "no loopback".
- the symbol ' $\star$ ', which means "all possible loopbacks from zero to length$l$ ". If no value is given the environment variable bmc_loopback is considered instead.

```
check_pslspec_sbmc [-h] [-m | -o output-file] [-n number | -p
"psl-expr [IN context]" | -P "name"] [-g] [-1] [-k
bmc_lenght] [-l loopback]
```

Check psl properties using SAT-based model checking. Use the SBMC algorithms.
A psl-expr to be checked can be specified at command line using option -p. Alternatively, option -n can be used for checking a particular formula in the property database. If neither -n nor -p are used, all the PSLSPEC formulas in the database are checked. Options -k and -1 can be used to define the maximum problem bound, and the value of the loopback for the single generated problems respectively; their values can be stored in the environment variables bmc_lenght and bmc_loopback. Single problems can be generated by using option -1 . Bounded model checking problems can be generated and dumped in a file by using option -g.
See variable use_coi_size_sorting for changing properties verification order.
Command Options:

## $-m$

- ○ output-file
-g
-l loopback
-p "psl-expr A PSL formula to be checked. context is the module instance name which [IN context]" the variables in psl-expr must be evaluated in.
$-n$ number Checks the PSL property with index number in the property database.
-P name $\quad$ Checks the PSL property named name in the property database.
-1 Generates a single bounded model checking problem with fixed bound and loopback values, it does not iterate incrementing the value of the problem bound.
-k bmc_length bmc_length is the maximum problem bound to be checked. Only natural numbers are valid values for this option. If no value is given the environment variable bmc_length is considered instead.
Pipes the output generated by the command in processing PSLSPECS to the program specified by the PAGER shell variable if defined, else through the UNIX command "more".
Writes the output generated by the command in processing PSLSPEC s to the file output-file

Dumps DIMACS version of bounded model checking problem into a file whose name depends on the system variable bmc_dimacs_filename. This feature is not allowed in combination of the option -i.

The loopback value may be:

- a natural number in ( 0 , max_length-1). A positive sign ('+') can be also used as prefix of the number. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- a negative number in (-1, -bmc_length). In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' $x$ ', which means "no loopback".
- the symbol ' $\star$ ', which means "all possible loopbacks from zero to length$l$ ". If no value is given the environment variable bmc_loopback is considered instead.

```
check_pslspec_sbmc_inc - Performs incremental SAT-based PSL model check-
ing
```

```
check_pslspec_sbmc_inc [-h] [-m | -o output-file] [-n number | -p
"psl-expr [IN context]" | -P "name"] [-1] [-k
bmc_lenght] [-l loopback] [-c] [-N]
```

Check psl properties using incremental SAT-based model checking. Use the SBMC algorithms.
A psl-expr to be checked can be specified at command line using option -p. Alternatively, option -n can be used for checking a particular formula in the property database. If neither -n nor -p are used, all the PSLSPEC formulas in the database are checked. Options -k and -1 can be used to define the maximum problem bound, and the value of the loopback for the single generated problems respectively; their values can be stored in the environment variables bmc_lenght and bmc_loopback. Single problems can be generated by using option -1 . Bounded model checking problems can be generated and dumped in a file by using option -g . With the option -c is possible to perform a completeness check, while with the option -N is possible to disable the virtual unrolling.

See variable use_coi_size_sorting for changing properties verification order.

## Command Options:

$$
\begin{aligned}
& -m \quad \text { Pipes the output generated by the command in processing PSLSPECS to the } \\
& \text { program specified by the PAGER shell variable if defined, else through the } \\
& \text { UNIX command "more". } \\
& \text {-○ output-file Writes the output generated by the command in processing PSLSPEC s to the } \\
& \text { file output-file } \\
& \text {-p "psl-expr A PSL formula to be checked. context is the module instance name which } \\
& \text { [IN context]" the variables in psl-expr must be evaluated in. } \\
& -\mathrm{n} \text { number } \quad \text { Checks the PSL property with index number in the property database. } \\
& -\mathrm{P} \text { name } \quad \text { Checks the PSL property named name in the property database. } \\
& \text {-1 } \\
& \text { - } \mathrm{k} \text { bmc_length bmc_length is the maximum problem bound to be checked. Only natural } \\
& \text { numbers are valid values for this option. If no value is given the environment } \\
& \text { variable bmc_length is considered instead. } \\
& \text {-l loopback } \\
& \text { The loopback value may be: } \\
& \text { - a natural number in ( } 0 \text {, max_length-1). A positive sign ('+') can be also } \\
& \text { used as prefix of the number. Any invalid combination of length and loop- } \\
& \text { back will be skipped during the generation/solving process. }
\end{aligned}
$$

- a negative number in (-1, -bmc_length). In this case loopback is considered a value relative to max_length. Any invalid combination of length and loopback will be skipped during the generation/solving process.
- the symbol ' $x$ ', which means "no loopback".
- the symbol ' $\star$ ', which means "all possible loopbacks from zero to length$l$ ". If no value is given the environment variable bmc_loopback is considered instead.
-c Performs completeness check.
$-\mathrm{N} \quad$ Does not perform virtual unrolling.


### 4.5 Simulation Commands

In this section we describe the commands that allow to simulate a NUXmV specification. See also the section Section 4.7 [Traces], page 88 that describes the commands available for manipulating traces.
pick_state - Picks a state from the set of initial states
Command
pick_state [-h] [-v] [-r | -i [-a]] [-c "constraints" | -s trace.state]
Chooses an element from the set of initial states, and makes it the current state (replacing the old one). The chosen state is stored as the first state of a new trace ready to be lengthened by steps states by the simulate command. The state can be chosen according to different policies which can be specified via command line options. By default the state is chosen in a deterministic way.

## Command Options:

| -v | Verbosely prints out chosen state (all state and frozen variables, otherwise it <br> prints out only the label $t .1$ of the state chosen, where $t$ is the number of <br> the new trace, that is the number of traces so far generated plus one). |
| :---: | :--- |
| -r | Randomly picks a state from the set of initial states. |
| -i | Enables the user to interactively pick up an initial state. The user is requested <br> to choose a state from a list of possible items (every item in the list doesn't <br> show frozen and state variables unchanged with respect to a previous item). <br> If the number of possible states is too high, then the user has to specify some <br> further constraints as "simple expression". <br> Displays all state and frozen variables (changed and unchanged with respect <br> to a previous item) in an interactive picking. This option works only if the <br> -i options has been specified. |
| -c | "constraints"Uses constraints to restrict the set of initial states in which the state has <br> to be picked. constraints must be enclosed between double quotes " ". <br> Picks state from trace.state label. A new simulation trace will be created by <br> copying prefix of the source trace up to specified state. |

simulate - Performs a simulation from the current selected state

```
simulate [-h] [-p | -v] [-r | -i [-a]] [-c "constraints" | -t
"constraints"] [-k steps]
```

Generates a sequence of at most steps states (representing a possible execution of the model), starting from the current state. The current state must be set via the pick_state or goto_state commands.

It is possible to run the simulation in three ways (according to different command line policies): deterministic (the default mode), random and interactive.
The resulting sequence is stored in a trace indexed with an integer number taking into account the total number of traces stored in the system. There is a different behavior in the way traces are built, according to how current state is set: current state is always put at the beginning of a new trace (so it will contain at most steps +1 states) except when it is the last state of an existent old trace. In this case the old trace is lengthened by at most steps states.

Command Options:
-p Prints current generated trace (only those variables whose value changed from the previous state).
$-\mathrm{v} \quad$ Verbosely prints current generated trace (changed and unchanged state and frozen variables).
$-r \quad$ Picks a state from a set of possible future states in a random way.
-i Enables the user to interactively choose every state of the trace, step by step. If the number of possible states is too high, then the user has to specify some constraints as simple expression. These constraints are used only for a single simulation step and are forgotten in the following ones. They are to be intended in an opposite way with respect to those constraints eventually entered with the pick_state command, or during an interactive simulation session (when the number of future states to be displayed is too high), that are local only to a single step of the simulation and are forgotten in the next one.
To improve readability of the list of the states which the user must pick one from, each state is presented in terms of difference with respect of the previous one.
-a Displays all the state and frozen variables (changed and unchanged) during every step of an interactive session. This option works only if the -i option has been specified.
-c "constraints" Performs a simulation in which computation is restricted to states satisfying those constraints. The desired sequence of states could not exist if such constraints were too strong or it may happen that at some point of the simulation a future state satisfying those constraints doesn't exist: in that case a trace with a number of states less than steps trace is obtained. Note: constraints must be enclosed between double quotes " ". The expression cannot contain next operators, and is automatically shifted by one state in order to constraint only the next steps
-t "constraints" Performs a simulation in which computation is restricted to states satisfying those constraints. The desired sequence of states could not exist if such constraints were too strong or it may happen that at some point of the simulation a future state satisfying those constraints doesn't exist: in that case a trace with a number of states less than steps trace is obtained. Note: constraints must be enclosed between double quotes " ". The expression can contain next operators, and is NOT automatically shifted by one state as done with option -c
-k steps Maximum length of the path according to the constraints. The length of a trace could contain less than steps states: this is the case in which simulation stops in an intermediate step because it may not exist any future state satisfying those constraints. The default value is determined by the default_simulation_steps environment variable

## default_simulation_steps

 Environment VariableControls the default number of steps performed by all simulation commands. The default is 10 .

## shown_states

Environment Variable
Controls the maximum number of states tail will be shown during an interactive simulation session. Possible values are integers from 1 to 100 . The default value is 25 .

## traces_hiding_prefix

Environment Variable
see section 4.7.2 for a detailed description.
traces_regexp
Environment Variable
see section 4.7.2 for a detailed description.

### 4.6 Execution Commands

In this section we describe the commands that allow to perform trace re-execution on a given model. See also the section Section 4.7 [Traces], page 88 that describes the commands available for manipulating traces.
execute_traces - Executes complete traces on the model FSM
Command

```
execute_traces [-h] [-v] [-m | -o output-file] -e engine [-a |
trace_number]
```

Executes traces stored in the Trace Manager. If no trace is specified, last registered trace is executed. Traces must be complete in order to perform execution.

Command Options:

| -v | Verbosely prints traces execution steps. |
| :--- | :--- |
| -a | Prints all the currently stored traces. |

$-m \quad$ Pipes the output through the program specified by the PAGER shell variable if defined, else through the UNIX command "more".
-o output-file Writes the output generated by the command to output-file.
-e engine Selects an engine for trace re-execution. It must be one of 'bdd', 'sat' or 'smt'.
trace_number
The (ordinal) identifier number of the trace to be printed. This must be the last argument of the command. Omitting the trace number causes the most recently generated trace to be executed.

```
execute_partial_traces [-h] [-v] [-r] [-m | -o output-file] -e engine
[-a | trace_number]
```

Executes traces stored in the Trace Manager. If no trace is specified, last registered trace is executed. Traces are not required to be complete. Upon succesful termination, a new complete trace is registered in the Trace Manager.

Command Options:

| -v | Verbosely prints traces execution steps. |
| :---: | :---: |
| -a | Prints all the currently stored traces. |
| -r | Performs restart on complete states. When a complete state (i.e. a state which is non-ambiguosly determined by a complete assignment to state variables) is encountered, the re-execution algorithm is re-initialized, thus reducing computation time. |
| -m | Pipes the output through the program specified by the PAGER shell variable if defined, else through the UNIX command "more". |
| -o output-file | Writes the output generated by the command to output-file. |
| -e engine | Selects an engine for trace re-execution. It must be one of 'bdd', 'sat' or 'smt'. |
| trace_number | The (ordinal) identifier number of the trace to be printed. This must be the last argument of the command. Omitting the trace number causes the most recently generated trace to be executed. |

### 4.7 Traces

A trace consists of an initial state, optionally followed by a sequence of states-inputs pairs corresponding to a possible execution of the model. Apart, from the initial state, each pair contains the inputs that caused the transition to the new state, and the new state itself. The initial state has no such input values defined as it does not depend on the values of any of the inputs. The values of any constants declared in DEFINE sections are also part of a trace. If the value of a constant depends only on state and frozen variables then it will be treated as if it is a state variable too. If it depends only on input variables then it will be treated as if it is an input variable. If however, a constant depends upon both input and state/frozen variables and/or NEXTed state variables, then it gets displayed in a separate "combinatorial" section. Since the values of any such constants depend on one or more inputs, the initial state does not contain this section either.

Traces are created by NUXMV when a formula is found to be false; they are also generated as a result of a simulation (Section 4.5 [Simulation Commands], page 85) or partial trace re-execution (Section 4.6 [Execution Commands], page 87). Each trace has a number, and the states-inputs pairs are numbered within the trace. Trace $n$ has states/inputs $n .1, n .2, n .3$, '...." where $n .1$ represents the initial state.

When Cone of Influence (COI) is enabled when generating a trace (e.g. when performing model checking), the generated trace will contain only the relevant symbols (variables and DEFINEs) which are in the COI projected by the variables occurring in the property which is being checked. The symbols which are left out of the COI, will be not visible in the generated trace, as they do not occur in the problem encoded in the solving engine. Notice that when COI is enabled, the generated trace may or may not be a valid counter-example trace for the original model.

### 4.7.1 Inspecting Traces

The trace inspection commands of NUXMV allow for navigation along the labelled states-inputs pairs of the traces produced. During the navigation, there is a current state, and the current trace is the trace the current state belongs to. The commands are the following:
goto_state - Goes to a given state of a trace

```
goto_state [-h] state_label
```

Makes state_label the current state. This command is used to navigate along traces produced by NuSMV. During the navigation, there is a current state, and the current trace is the trace the current state belongs to.
state_label is in the form trace.state where
trace is the index of the trace which the state has to be taken from.
state is the index of the state within the given trace. If state is a negative number, then the state index is intended to be relative to the length of the given trace. For example 2.-1 means the last state of the trace 2. 2.-2 is the state before the last state, etc.

## print_current_state - Prints out the current state

```
print_current_state [-h] [-v]
```

Prints the name of the current state if defined.
Command Options:
$-\mathrm{v} \quad$ Prints the value of all the state and frozen variables of the current state.

### 4.7.2 Displaying Traces

nUXMV comes with three trace plugins (see Section 4.8 [Trace Plugins], page 92) which can be used to display traces in the system. Once a trace has been generated by NUXMV it is printed to stdout using the trace explanation plugin which has been set as the current default. The command show_traces (see Section 4.5 [Simulation Commands], page 85) can then be used to print out one or more traces using a different trace plugin, as well as allowing for output to a file.

Generation and displaying of traces can be enabled/disabled by setting variable counter_examples. Some filtering of symbols that are presented when showing traces can be controlled by variables traces_hiding_prefix and traces_regexp.
counter_examples Environment Variable

This determines whether traces are generated when needed. See also command line option -dcx.

Symbols names that match this string prefix will be not printed out when showing a trace. This variable may be used to avoid displaying symbols that are expected to be not visible to the user. For example, this variable is exploited when dumping booleanized models, as NUXMV may introduce hidden placeholding symbols as DEFINES that do not carry any useful information for the user, and that would make traces hardly readable if printed. Default is __

Only symbols whose names match this regular expression will be printed out when showing a trace. This option might be used by users that are interested in showing only some symbol names. Names are first filtered out by applying matching of the dual variable traces_hiding_prefix, and then filtered names are checked against content of traces_regexp. Given regular expression can be a Posix Basic Regular Expression. Matching is carried out on symbol names without any contextual information, like module hierarchy. For example in $\mathrm{m} 1 . \mathrm{m} 2$. name only name is checked for filtering.
Notice that depending on the underlaying platform and operating system this variable might be not available.

Controls whether defines should be printed as part of a trace or be skipped. Skipping printing of the defines can help in reducing time and memory usage required to build very big traces.

Controls whether defines containing next operators should be printed as part of a trace or be skipped.

### 4.7.3 Trace Plugin Commands

The following commands relate to the plugins which are available in NUXMV.
show_plugins - Shows the available trace explanation plugins

```
show_plugins [-h] [-n plugin-no | -a]
```

Command Options:

```
-n plugin-no Shows the plugin with the index number equal to plugin-no.
-a Shows all the available plugins.
```

Shows the available plugins that can be used to display a trace which has been generated by NUSMV, or that has been loaded with the read_trace command. The plugin that is used to read in a trace is also shown. The current default plugin is marked with " $[D]$ ".
All the available plugins are displayed by default if no command options are given.

## default_trace_plugin

Environment Variable
This determines which trace plugin will be used by default when traces that are generated by NUXMV are to be shown. The values that this variable can take depend on which trace plugins are installed. Use the command show_plugins to see which ones are available. The default value is 0 .

```
show_traces [-h] [-v] [-t] [-A] [-m | -o output-file]
[-p plugin-no] [-a | trace_number[.from_state[:[to_state]]]
```

Command Options:

| -v | Verbosely prints traces content (all state and frozen variables, otherwise it prints out only those variables that have changed their value from previous state). This option only applies when the Basic Trace Explainer plugin is used to display the trace. |
| :---: | :---: |
| -t | Prints only the total number of currently stored traces. |
| -a | Prints all the currently stored traces. |
| -m | Pipes the output through the program specified by the PAGER shell variable if defined, else through the UNIX command "more". |
| -o output-file | Writes the output generated by the command to output-file. |
| -p plugin-no | Uses the specified trace plugin to display the trace. |
| trace_number | The (ordinal) identifier number of the trace to be printed. Omitting the trace number causes the most recently generated trace to be printed. |
| from_step | The number of the first step of the trace to be printed. Negative numbers can be used to denote right-to-left indexes from the last step. |
| to_step | The number of the trace to be printed. Negative numbers can be used to denote right-to-left indexes from the last step. Omitting this parameter causes the entire suffix of the trace to be printed. |
| -A | Prints the trace(s) using a rewriting mapping for all symbols. The rewriting is the same used in write_flat model with option -A. |

Verbosely prints traces content (all state and frozen variables, otherwise it prints out only those variables that have changed their value from previous state). This option only applies when the Basic Trace Explainer plugin is used to display the trace.
-t Prints only the total number of currently stored traces.
Prints all the currently stored traces.
Pipes the output through the program specified by the PAGER shell variable if defined, else through the UNIX command "more".

Writes the output generated by the command to output-file.
Uses the specified trace plugin to display the trace. number causes the most recently generated trace to be printed. be used to denote right-to-left indexes from the last step.

Shows the traces currently stored in system memory, if any. By default it shows the last generated trace, if any. Optional trace number can be followed by two indexes (from_state, to_state), denoting a trace "slice". Thus, it is possible to require printout only of an arbitrary fragment of the trace (this can be helpful when inspecting very big traces).

If the XML Format Output plugin is being used to save generated traces to a file with the intent of reading them back in again at a later date, then only one trace should be saved per file. This is because the trace reader does not currently support multiple traces in one file.

```
read_trace [-h | [-i filename] [-u] [-s] filename]
```


## Command Options:

-i filename Reads in a trace from the specified file. Note that the file must only contain one trace. This option has been deprecated. Use the explicit filename argument instead.
-u Turns "undefined symbol" error into a warning. The loader will ignore assignments to undefined symbols.
-s Turns "wrong section" error into a warning. The loader will accept symbol assignment even if they are in a different section than expected. Assignments will be silently moved to appropriate section, i.e. misplaced assignments to state symbols will be moved back to previous state section and assignments to input/combinatorial symbols will be moved forward to successive input/combinatorial section. Such a way if a variable in a model was input and became state or vice versa the existing traces still can be read and executed.

Loads a trace which has been previously output to a file with the XML Format Output plugin. The model from which the trace was originally generated must be loaded and built using the command "go" first.
Please note that this command is only available on systems that have the libxml2 XML parser library installed.

### 4.8 Trace Plugins

nUXmV comes with three plugins which can be used to display a trace that has been generated:

Basic Trace Explainer<br>States/Variables Table<br>XML Format Printer

There is also an xml loader which can read in any trace which has been output to a file by the XML Format Printer. Note however that this loader is only available on systems that have the libxml2 XML parser library installed.

Once a trace has been generated it is output to stdout using the currently selected plugin. The command show_traces can be used to output any previuosly generated, or loaded, trace to a specific file.

### 4.8.1 Basic Trace Explainer

This plugin prints out each state (the current values of the variables) in the trace, one after the other. The initial state contains all the state and frozen variables and their initial values. States are numbered in the following fashion:

```
trace_number.state_number
```

There is the option of printing out the value of every variable in each state, or just those which have changed from the previous one. The one that is used can be chosen by selecting the appropriate trace plugin. The values of any constants which depend on both input and state or frozen variables are printed next. It then prints the set of inputs which cause the transition to a new state (if the model contains inputs), before actually printing the new state itself. The set of inputs and the subsequent state have the same number associated to them.

In the case of a looping trace, if the next state to be printed is the same as the last state in the trace, a line is printed stating that this is the point where the loop begins.

With the exception of the initial state, for which no input values are printed, the output syntax for each state is as follows:

```
-> Input: TRACE_NO.STATE_NO <-
    /* for each input var (being printed), i: */
    INPUT_VARi = VALUE
-> State: TRACE_NO.STATE_NO <-
    /* for each state and frozen var (being printed), j: */
    STATE_VARj = VALUE
    /* for each combinatorial constant (being printed), k: */
    CONSTANTk = VALUE
```

where INPUT_VAR, STATE_VAR and CONSTANT have the relevant module names prepended to them (seperated by a period) with the exception of the module "main".

The version of this plugin which only prints out those variables whose values have changed is the initial default plugin used by nuXmv.

### 4.8.2 States/Variables Table

This trace plugin prints out the trace as a table, either with the states on each row, or in each column. The entries along the state axis are:
S1 C2 I2 S2 ... Cn In Sn
where S1 is the initial state, and $I_{i}$ gives the values of the input variables which caused the transition from state $S_{i-1}$ to state $S_{i} . C_{i}$ gives the values of any combinatorial constants, where the value depends on the values of the state or frozen variables in state $S_{i-1}$ and the values of input variables in state $S_{i}$.

The variables in the model are placed along the other axis. Only the values of state and frozen variables are displayed in the State row/column, only the values of input variables are displayed in the Input row/column and only the values of combinatorial constants are displayed in the Constants row/column. All remaining cells have '-' displayed.

### 4.8.3 XML Format Printer

This plugin prints out the trace either to stdout or to a specified file using the command show_traces. If traces are to be output to a file with the intention of them being loaded again at a later date, then each trace must be saved in a separate file. This is because the XML Reader plugin does not currently support multiple traces per file.
The format of a dumped XML trace file is as follows:

```
<?XML_VERSION_STRING?>
<counter-example type=TRACE_TYPE desc=TRACE_DESC>
    /* for each state, i: */
    <node>
        <state id=i>
            /* for each state and frozen var, j: */
            <value variable=j>VALUE</value>
        </state>
        <combinatorial id=i+1>
            /* for each combinatorial constant, k: */
            <value variable=k>VALUE</value>
        </combinatorial>
        <input id=i+1>
            /* for each input var, l: */
            <value variable=l>VALUE</value>
        </input>
    </node>
</counter-example>
```

Note that for the last state in the trace, there is no input section in the node tags. This is because the inputs section gives the new input values which cause the transition to the next state in the trace. There is also no combinatorial section as this depends on the values of the inputs and are therefore undefined when there are no inputs.

### 4.8.4 XML Format Reader

This plugin makes use of the libxml2 XML parser library and as such can only be used on systems where this library is available. Previously generated traces for a given model can be loaded using this plugin provided that the original model file ${ }^{1}$ has been loaded, and built using the command $g o$.

When a trace is loaded, it is given the smallest available trace number to identify it. It can then be manipulated in the same way as any generated trace.

### 4.9 Interface to the DD Package

nUXMV uses the state of the art BDD package CUDD [Som98]. Control over the BDD package can be very important to tune the performance of the system. In particular, the order of variables is critical to control the memory and the time required by operations over BDDs. Reordering methods can be activated to determine better variable orders, in order to reduce the size of the existing BDDs.

Reordering of the variables can be triggered in two ways: by the user, or by the BDD package. In the first way, reordering is triggered by the interactive shell command dynamic_var_ordering with the $-f$ option.

Reordering is triggered by the BDD package when the number of nodes reaches a given threshold. The threshold is initialized and automatically adjusted after each reordering by the package. This is called dynamic reordering, and can be enabled or disabled by the user. Dynamic reordering is enabled with the shell command dynamic_var_ordering with the option -e, and disabled with the - doption. Variable dynamic_reorder can also be used to determine whether dynamic reordering is active. If dynamic reordering is enabled it may be beneficial also to disable BDD caching by unsetting variable enable_sexp2bdd_caching.

## dynamic_reorder <br> Environment Variable

Determines whether dynamic reordering is active. If this variable is set, dynamic reordering will take place as described above. If not set (default), no dynamic reordering will occur. This variable can also be set by passing -dynamic command line option when invoking nUXmV.
reorder_method Environment Variable

Specifies the ordering method to be used when dynamic variable reordering is fired. The possible values, corresponding to the reordering methods available with the CUDD package, are listed below. The default value is sift.
sift:
random:
random_pivot:

Moves each variable throughout the order to find an optimal position for that variable (assuming all other variables are fixed). This generally achieves greater size reductions than the window method, but is slower.

Pairs of variables are randomly chosen, and swapped in the order. The swap is performed by a series of swaps of adjacent variables. The best order among those obtained by the series of swaps is retained. The number of pairs chosen for swapping equals the number of variables in the diagram.
Same as random, but the two variables are chosen so that the first is above the variable with the largest number of nodes, and the second is below that variable. In case there are several variables tied for the maximum number of nodes, the one closest to the root is used.

[^9]```
sift_converge:
symmetry_sift:
symmetry_sift_converge:
window2:
window3:
window4:
window2_converge:
window3_converge:
window4_converge:
group_sift:
group_sift_converge:
annealing:
```

genetic: This method is an implementation of a genetic algo-
rithm for variable ordering. This method is poten-
tially very slow.

This method implements a dynamic programming approach to exact reordering. It only stores one BDD at a time. Therefore, it is relatively efficient in terms of memory. Compared to other reordering strategies, it is very slow, and is not recommended for more than 16 boolean variables.
This method is a combination of sifting and linear transformations.
linear_conv: The linear method is iterated until no further improvement is obtained.

```
dynamic_var_ordering [-d] [-e <method>] [-f <method>] [-h]
```

Controls the application and the modalities of (dynamic) variable ordering. Dynamic ordering is a technique to reorder the BDD variables to reduce the size of the existing BDDs. When no options are specified, the current status of dynamic ordering is displayed. At most one of the options $-e,-f$, and -d should be specified. Dynamic ordering may be time consuming, but can often reduce the size of the BDDs dramatically. A good point to invoke dynamic ordering explicitly (using the $-f$ option) is after the commands build_model, once the transition relation has been built. It is possible to save the ordering found using write_order in order to reuse it (using build_model -i order-file) in the future.

Command Options

```
-d
-e <method>
```

-f <method>

Disable dynamic ordering from triggering automatically.
Enable dynamic ordering to trigger automatically whenever a certain threshold on the overall BDD size is reached. <method> must be one of the following:

- sift: Moves each variable throughout the order to find an optimal position for that variable (assuming all other variables are fixed). This generally achieves greater size reductions than the window method, but is slower.
- random: Pairs of variables are randomly chosen, and swapped in the order. The swap is performed by a series of swaps of adjacent variables. The best order among those obtained by the series of swaps is retained. The number of pairs chosen for swapping equals the number of variables in the diagram.
- random_pivot: Same as random, but the two variables are chosen so that the first is above the variable with the largest number of nodes, and the second is below that variable. In case there are several variables tied for the maximum number of nodes, the one closest to the root is used.
- sift_converge: The sift method is iterated until no further improvement is obtained.
- symmetry_sift: This method is an implementation of symmetric sifting. It is similar to sifting, with one addition: Variables that become adjacent during sifting are tested for symmetry. If they are symmetric, they are linked in a group. Sifting then continues with a group being moved, instead of a single variable.
- symmetry_sift_converge: The symmetry_sift method is iterated until no further improvement is obtained.
- window $\{\mathbf{2}, \mathbf{3}, \mathbf{4}\}$ : Permutes the variables within windows of "n" adjacent variables, where " n " can be either 2,3 or 4 , so as to minimize the overall BDD size.
- window $\{\mathbf{2}, \mathbf{3}, \mathbf{4}\}$ _converge: The window $\{\mathbf{2}, \mathbf{3}, \mathbf{4}\}$ method is iterated until no further improvement is obtained.
- group_sift: This method is similar to symmetry_sift, but uses more general criteria to create groups.
- group_sift_converge: The group_sift method is iterated until no further improvement is obtained.
- annealing: This method is an implementation of simulated annealing for variable ordering. This method is potentially very slow.
- genetic: This method is an implementation of a genetic algorithm for variable ordering. This method is potentially very slow.
- exact: This method implements a dynamic programming approach to exact reordering. It only stores a BDD at a time. Therefore, it is relatively efficient in terms of memory. Compared to other reordering strategies, it is very slow, and is not recommended for more than 16 boolean variables.
- linear: This method is a combination of sifting and linear transformations.
- linear_converge: The linear method is iterated until no further improvement is obtained.
Force dynamic ordering to be invoked immediately. The values for <method> are the same as in option -e.

```
clean_sexp2bdd_cache [-h]
```

During conversion of symbolic expressions to ADD and BDD representations the results of evaluations are normally cached (see additionally the environment variable enable_sexp2bdd_caching). This allows to save time by avoid the construction of BDD for the same symbolic expression several time.
In some situations it may be preferable to clean the cache and free collected ADD and BDD. This operation can be done, for example, to free some memory. Another possible reason is that dynamic reordering may modify all existing BDDs, and cleaning the cache thereby freeing the BDD may speed up the reordering.

This command is designed specifically to free the internal cache of evaluated expressions and their ADDs and BDDs. Note that only the cache of symbolic-expression-to-bdd evaluator is freed. BDDs of variables, constants and expressions collected in BDD FSM or anywhere else are not freed.
print_formula - Prints a formula in canonical format.
Command
print_formula [-h] [-v] [-f] "expression"
Prints the number of satsfying assignments for the given formula. In verbose mode, prints also the list of such assigments. In formula mode, a canonical representation of the formula is printed.

Command Options:

$$
\begin{array}{ll}
-\mathrm{v} & \text { Prints explicit models of the formula. } \\
-\mathrm{f} & \text { Prints the simplified and canonical formula. }
\end{array}
$$

This variable determines if during evaluation of symbolic expression to ADD and BDD representations the obtained results are cached or not. Note that if the variable is set down consequently computed results are not cached but the previously cached data remain unmodified and will be used during later evaluations.
The default value of this variable is 1 which can be changed by a command line option -disable_sexp2bdd_caching.
For more information about the reasons of why BDD cache should be disabled in some situations see command clean_sexp2bdd_cache.
print_bdd_stats - Prints out the BDD statistics and parameters
Command

```
print_bdd_stats [-h]
```

Prints the statistics for the BDD package. The amount of information depends on the BDD package configuration established at compilation time. The configurtion parameters are printed out too. More information about statistics and parameters can be found in the documentation of the CUDD Decision Diagram package.
set_bdd_parameters - Creates a table with the value of all currently active
Command
NuSMV flags and change accordingly the configurable parameters of the BDD package.

```
set_bdd_parameters [-h] [-s]
```

Applies the variables table of the NUSMV environnement to the BDD package, so the user can set specific BDD parameters to the given value. This command works in conjunction with the print_bdd_stats and set commands. print_bdd_stats first prints a report of the parameters and statistics of the current bdd_manager. By using the command set, the user may modify the value of any of the parameters of the underlying BDD package. The way to do it is by setting a value in the variable BDD. parameter name where parameter name is the name of the parameter exactly as printed by the print_bdd_stats command.


## Command Options:

$-s \quad$ Prints the BDD parameter and statistics after the modification.

### 4.10 Administration Commands

This section describes the administrative commands offered by the interactive shell of NUXMV.
!- shell_command Command
"! " executes a shell command. The "shell_command" is executed by calling "bin/sh -c shell_command". If the command does not exists or you have not the right to execute it, then an error message is printed.
alias [-h] [<name> [<string>]]
The alias command, if given no arguments, will print the definition of all current aliases. Given a single argument, it will print the definition of that alias (if any). Given two arguments, the keyword <name> becomes an alias for the command string <string>, replacing any other alias with the same name.

## Command Options:

```
<name>
Alias
<string> Command string
```

It is possible to create aliases that take arguments by using the history substitution mechanism. To protect the history substitution character ' $\%$ ' from immediate expansion, it must be preceded by a ' $\backslash$ ' when entering the alias.

For example:

```
nuXmv > alias read "read_model -i %:1.smv ; set input_order_file %:1.ord"
```

nuXmv > read short
will create an alias 'read', execute "read_model -i short.smv; set input_order_file short.ord". And again:
nuXmv > alias echo2 "echo Hi ; echo \%* !"
nuXmv > echo2 happy birthday
will print:
Hi
happy birthday !

CAVEAT: Currently there is no check to see if there is a circular dependency in the alias definition. e.g.
nuXmv > alias foo "echo print_bdd_stats; foo"
creates an alias which refers to itself. Executing the command foo will result an infinite loop during which the command print_bdd_stats will be executed.
echo [-h] [-2] [-n] [-o filename [-a]] <string>
Echoes the specified string either to standard output, or to filename if the option $-\circ$ is specified.

## Command Options:

```
-2
-n
-o filename
```

-a

Redirects output to the standard error instead of the standard output. This cannot be used in combination with the option -o.
Does not output the trailing newline.
Echoes to the specified filename instead of to standard output. If the option -a is not specified, the file filename will be overwritten if it already exists.
Appends the output to the file specified by option $-\bigcirc$, instead of overwritting it. Use only with the option -o.

```
help [-a] [-h] [<command>]
```

If invoked with no arguments help prints the list of all commands known to the command interpreter. If a command name is given, detailed information for that command will be provided.

Command Options:
-a Provides a list of all internal commands, whose names begin with the underscore character (' ${ }^{\prime}$ ) by convention.

## history - list previous commands and their event numbers

```
history [-h] [<num>]
```

Lists previous commands and their event numbers. This is a UNIX-like history mechanism inside the NuSMV shell.

## Command Options:

<num> Lists the last <num> events. Lists the last 30 events if <num> is not specified.

## History Substitution:

The history substitution mechanism is a simpler version of the csh history substitution mechanism. It enables you to reuse words from previously typed commands.

The default history substitution character is the ' $\%$ ' ('!’ is default for shell escapes, and ' $\#$ ' marks the beginning of a comment). This can be changed using the set command. In this description ' $\%$ ' is used as the history_char. The ' $\%$ ' can appear anywhere in a line. A line containing a history substitution is echoed to the screen after the substitution takes place. ' $\%$ ' can be preceded by a 'ín order to escape the substitution, for example, to enter a ' $\%$ ' into an alias or to set the prompt.
Each valid line typed at the prompt is saved. If the history variable is set (see help page for set), each line is also echoed to the history file. You can use the history command to list the previously typed commands.

Substitutions:
At any point in a line these history substitutions are available.
Command Options:

Fovidione

\begin{tabular}{ll}
$\%: 0$ \& Initial word of last command. <br>
$\%: n$ \& n -th argument of last command. <br>
$\% \$$ \& Last argument of last command. <br>
$\% \star$ \& All but initial word of last command. <br>
$\% \%$ \& Last command. <br>
$\%$ stuf \& Last command beginning with "stuf". <br>
$\% \mathrm{n}$ \& Repeat the n-th command. <br>
$\%-\mathrm{n}$ \& Repeat the n-th previous command. <br>

old`new \& | Replace "old" with "new" in previous command. Trailing spaces are signif- |
| :--- |
| icant during substitution. Initial spaces are not significant. |

\end{tabular}

print_usage - Prints processor and BDD statistics.

```
print_usage [-h]
```

Prints a formatted dump of processor-specific usage statistics, and BDD usage statistics. For Berkeley Unix, this includes all of the information in the getrusage () structure.

```
quit - exits NuSMV
```

```
quit [-h] [-s] [-x]
```

Stops the program. Does not save the current network before exiting.

Command Options:

| $-s$ | Frees all the used memory before quitting. This is slower, and it is used for <br> finding memory leaks. |
| :--- | :--- |
| -x | Leaves immediately. Skip all the cleanup code, leaving it to the OS. This <br> can save quite a long time. |

reset - Resets the whole system.
Command
reset [-h]
Resets the whole system, in order to read in another model and to perform verification on it.
set - Sets an environment variable
Command

```
set [-h] [<name>] [<value>]
```

A variable environment is maintained by the command interpreter. The set command sets a variable to a particular value, and the unset command removes the definition of a variable. If set is given no arguments, it prints the current value of all variables.

## Command Options:

<name> Variable name
<value> Value to be assigned to the variable.

Using the set command to set a variable, without giving any explicit value is allowed, and sets the variable to 1 :
nuXmv > set foo
will set the variable foo to 1 .

Interpolation of variables is allowed when using the set command. The variables are referred to with the prefix of '\$'. So for example, what follows can be done to check the value of a set variable:
nuXmv > set foo bar
nuXmv > echo \$foo
bar

The last line "bar" will be the output produced by NUSMV. Variables can be extended by using the character ' $:$ ' to concatenate values. For example:
nuXmv > set foo bar
nuXmv > set foo \$foo:foobar
nuXmv > echo \$foo
bar:foobar

The variable foo is extended with the value foobar. Whitespace characters may be present within quotes. However, variable interpolation lays the restriction that the characters ' $\because$ ' and ' $l$ ' may not be used within quotes. This is to allow for recursive interpolation. So for example, the following is allowed

```
nuXmv > set "foo bar" this
nuXmv > echo $"foo bar"
this
```

The last line will be the output produced by NuSMV.
But in the following, the value of the variable foo/bar will not be interpreted correctly: nuXmv > set
"foo/bar" this
nuXmv > echo \$"foo/bar"
foo/bar

If a variable is not set by the set command, then the variable is returned unchanged. Different commands use environment information for different purposes. The command interpreter makes use of the following parameters:

## Command Options:

| autoexec | Defines a command string to be automatically executed after every command <br> processed by the command interpreter. This is useful for things like timing <br> commands, or tracing the progress of optimization. |
| :--- | :--- |
| open_path | "open_path" (in analogy to the shell-variable PATH) is a list of colon- <br> separated strings giving directories to be searched whenever a file is opened <br> for read. Typically the current directory (.) is the first item in this list. The |
| standard system library (typically NUXMV_LIBRARY_PATH) is always im- |  |
| plicitly appended to the current path. This provides a convenient short-hand |  |
| mechanism for reaching standard library files. |  |

nusmv_stderr Standard error (normally ( stderr)) can be re-directed to a file by setting the variable nusmv_stderr.
nusmv_stdout Standard output (normally ( stdout)) can be re-directed to a file by setting the variable nusmv_stdout.
source [-h] [-p] [-s] [-x] <file> [<args>]
Reads and executes commands from a file.
Command Options:

| -p | Prints a prompt before reading each command. |
| :--- | :--- |
| -s | Silently ignores an attempt to execute commands from a nonexistent file. |
| -x | Echoes each command before it is executed. |
| $\langle\mathrm{fil} \mathrm{l}\rangle$ | File name. |

Arguments on the command line after the filename are remembered but not evaluated. Commands in the script file can then refer to these arguments using the history substitution mechanism. EXAMPLE: Contents of test.scr:

```
read_model -i %:2
flatten_hierarchy
build_variables
build_model
compute_fairness
```

Typing source test.scr short.smv on the command line will execute the sequence

```
read.model -i short.smv
flatten_hierarchy
build_variables
build_model
compute_fairness
```

(In this case $\%: 0$ gets source, $\%: 1$ gets test.scr, and $\%: 2$ gets short.smv.) If you type
alias st source test.scr and then type st short.smv bozo, you will execute
read_model -i bozo
flatten_hierarchy
build_variables
build_model
compute_fairness
because bozo was the second argument on the last command line typed. In other words, command substitution in a script file depends on how the script file was invoked. Switches passed to a command are also counted as positional parameters. Therefore, if you type st -x short. smv bozo, you will execute
$\underset{\substack{\text { FONDOAIONE } \\ \text { BRINOK KESS ER }}}{ }$

```
read_model -i short.smv
flatten_hierarchy
build_variables
build_model
compute_fairness
```

To pass the -x switch (or any other switch) to source when the script uses positional parameters, you may define an alias. For instance, alias srcx source -x .
See the variable on_failure_script_quits for further information.

```
time [-h]
```

Prints the processor time used since the last invocation of the time command, and the total processor time used since NUSMV was started.

```
unalias [-h] <alias-names>
```

Removes the definition of an alias specified via the alias command.

## Command Options:

> <alias-names> Aliases to be removed
$\underset{\substack{\text { FoNDOZZOOE } \\ \text { BRUNO KESER }}}{ }$
unset - Unsets an environment variable

```
unset [-h] <variables>
```

A variable environment is maintained by the command interpreter. The set command sets a variable to a particular value, and the unset command removes the definition of a variable.

## Command Options:

<variables> Variables to be unset.

## usage - Provides a dump of process statistics

```
usage [-h]
```

Prints a formatted dump of processor-specific usage statistics. For Berkeley Unix, this includes all of the information in the getrusage() structure.
which - Looks for a file called "file_name"

```
which [-h] <file_name>
```

Looks for a file in a set of directories which includes the current directory as well as those in the NUSMV path. If it finds the specified file, it reports the found file's path. The searching path is specified through the set open_path command in . nusmvrc.

## Command Options:

<file_name> File to be searched

### 4.11 Other Environment Variables

The behavior of the system depends on the value of some environment variables. For instance, an environment variable specifies the partitioning method to be used in building the transition relation. The value of environment variables can be inspected and modified with the "set" command. Environment variables can be either logical or utility.

Defines a command string to be automatically executed after every command processed by the command interpreter. This may be useful for timing commands, or tracing the progress of optimization.

## on_failure_script_quits

When a non-fatal error occurs during the interactive mode, the interactive interpreter simply stops the currently executed command, prints the reason of the problem, and prompts for a new command. When set, this variables makes the command interpreter quit when an error occur, and then quit nUXmV. This behaviour might be useful when the command source is controlled by either a system pipe or a shell script. Under these conditions a mistake within the script interpreted by source or any unexpected error might hang the controlling script or pipe, as by default the interpreter would simply give up the current execution, and wait for further commands. The default value of this environment variable is 0 .

Enables file completion a la "csh". If the system has been compiled with the "readline" library, the user is able to perform file completion by typing the $\langle\mathrm{TAB}\rangle$ key (in a way similar to the file completion inside the "bash" shell). If the system has not been compiled with the "readline" library, a built-in method to perform file completion a la "csh" can be used. This method is enabled with the 'set filec' command. The "csh" file completion method can be also enabled if the "readline" library has been used. In this case the features offered by "readline" will be disabled.

## shell_char <br> Environment Variable

shell_char specifies a character to be used as shell escape. The default value of this environment variable is '!'.
history_char Environment Variable
history_char specifies a character to be used in history substitutions. The default value of this environment variable is ' $\%$ '.

Environment Variable
open_path (in analogy to the shell-variable PATH) is a list of colon-separated strings giving directories to be searched whenever a file is opened for read. Typically the current directory (.) is first in this list. The standard system library (NUXMV_LIBRARY_PATH) is always implicitly appended to the current path. This provides a convenient short-hand mechanism for reaching standard library files.

Standard error (normally stderr) can be re-directed to a file by setting the variable nusmv_stderr.
nusmv_stdout Environment Variable

Standard output (normally stdout) can be re-directed to a file by setting the internal variable nusmv_stdout.

Standard input (normally stdin) can be re-directed to a file by setting the internal variable nusmv_stdin.

## Chapter 5

## Commands of NUXMV

In the following we present the new commands provided by nUXMV. Similarly to the case of the commands inherited from NUSMV, we also describe the environment variables that may affect the behavior of the commands. All the commands have been classified in different categories.

### 5.1 Commands for Model Simulation

In this section we describe the new commands that allow to simulate a NUXMV specification that may contains Integers and Reals. See also the section Section 4.7 [Traces], page 88 that describes the commands available for manipulating traces.
msat_pick_state - Picks a state from the set of initial states
msat_pick_state [-h] [-v] [-i [-a]] [-c "constraint" | -s trace.state]
Chooses an element from the set of initial states, and makes it the current state (replacing the old one). The chosen state is stored as the first state of a new trace ready to be lengthened by steps states by the simulate command. The state can be chosen according to different policies which can be specified via command line options. By default the state is chosen in a deterministic way.

Command Options:

$$
\begin{array}{ll}
\text {-v } & \begin{array}{l}
\text { Verbosely prints out chosen state (all state and frozen variables, otherwise it } \\
\text { prints out only the label } t .1 \text { of the state chosen, where } t \text { is the number of } \\
\text { the new trace, that is the number of traces so far generated plus one). }
\end{array} \\
\text { Enables the user to interactively pick up an initial state. The user is requested } \\
\text { to choose a state from a list of possible items (every item in the list doesn't } \\
\text { show frozen and state variables unchanged with respect to a previous item). } \\
\text { If the number of possible states is too high, then the user has to specify some } \\
\text { further constraints as "simple expression". } \\
\text { Displays all state and frozen variables (changed and unchanged with respect } \\
\text { to a previous item) in an interactive picking. This option works only if the } \\
\text {-i options has been specified. }
\end{array}
$$

-s trace.state Picks state from trace.state label. A new simulation trace will be created by copying prefix of the source trace up to specified state.

```
msat_simulate [-h] [-v] [-i [-a]] [-e] [-l] [-k length] [[-c
"simple_expr"] | [-t "next_expr"] | [-p "formula"]]
```

msat_simulate does not require a specification to build the problem, because only the model is used to build it. The problem length is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length.

Command Options:
-V
-e
-1
-i
-c simple_expr
-c simple_expr
-t next_expr
-p "formula" Performs a simulation in which computation is restricted to states satisfying the given LTL formula. Option -e cannot be used in conjunction with this option.
-k length Maximum length of the path according to the constraints. The length of a trace could contain less than length states: this is the case in which simulation stops in an intermediate step because it may not exist any future state satisfying those constraints. The default value is determined by the default_simulation_steps environment variable

```
bmc_inc_simulate [-h] [-p | -v] [-r | -i [-a]] [[-c "constraints"] | [-t
"constraints"] ] [-k steps]
```

Performs incremental simulation of the model. bmc_inc_simulate does not require a specification to build the problem, because only the model is used to build it. The problem length is represented by the -k command parameter, or by its default value stored in the environment variable bmc_length.

Command Options:

| -p | Prints the generated trace (only changed variables). |
| :--- | :--- |
| -v | Prints the generated trace (all variables). |
| -r | Picks a state from a set of possible future states in a random way. |
| -i | Enters simulation's interactive mode. |
| -a | Displays all the state variables (changed and unchanged) in the interactive <br> session |
| -c constraint | Performs a simulation in which computation is restricted to states satisfy- <br> ing those constraints. The desired sequence of states could not exist if <br> such constraints were too strong or it may happen that at some point of the <br> simulation a future state satisfying those constraints doesn't exist: in that <br> case a trace with a number of states less than steps trace is obtained. Note: <br> constraints must be enclosed between double quotes " ". The expres- <br> sion cannot contain next operators, and is automatically shifted by one state <br> in order to constraint only the next steps |

-t "constraints" Performs a simulation in which computation is restricted to states satisfying those constraints. The desired sequence of states could not exist if such constraints were too strong or it may happen that at some point of the simulation a future state satisfying those constraints doesn't exist: in that case a trace with a number of states less than steps trace is obtained. Note: constraints must be enclosed between double quotes " ". The expression can contain next operators, and is NOT automatically shifted by one state as done with option -c
-k steps Maximum length of the path according to the constraints. The length of a trace could contain less than steps states: this is the case in which simulation stops in an intermediate step because it may not exist any future state satisfying those constraints. The default value is determined by the default_simulation_steps environment variable

### 5.2 Commands for Invariant Checking

In this section we list the new commands for checking invariants. Some of these commands can be applied only to finite-state models (e.g. "check_invar_guided"), some others can be applied both to finite and infinite-state models (e.g. "check_invar_ic3").

```
check_invar_guided [-n index | -p "prop" | -P "name"] [-s] [-S] [-R]
[-d] [-a] [-u] [-f] (-h | -e "sere_expr" | -i sere_file)
```

Performs invariant checking using the given strategy on current model. Checking the invariant is the process of determining that all states reachable from the initial states of the model lie in the invariant. By default the computation of reachable states is done in two steps. At first, states satisfying the strategy are computed until fixpoint is reached. Then the forward image is applied to the computed states independently of the strategy until all the reachable states are detected.

Invariants to be verified have to be provided as simple formulas (the only temporal operator allowed is "next") in the input file via the INVARSPEC keyword or directly at command line, using the option -p .
Option $-n$ and $-P$ can be used for checking a particular invariant of the model. If neither $-n$ nor -p nor -P are used, all the invariants are checked.
The strategy must be a valid PSL formula. Allowed PSL operators are: "; $[*][* N]($ with $N>0) \mid "$

## Command Options:

```
-p "invar-expr The command line specified invariant formula to be verified. context is
    [IN context]" the module instance name which the variables in invar-expr must be
                    evaluated in.
-n "index" Verifies the invariant with index "index" within the Property Database
-P "name" Verifies the invariant named "name" within the Property Database
-s Uses model simplification over the given model
-S Enables the use of a further simplified FSM for each atomic part of the given
    SERE
-R Enables the use of Implicit Frame Condition for each atomic part of the given
    SERE
-d Disables the reachability analysis completion. This means that only the strat-
    egy provided with the command is executed. The resulting set of reachable
    states is not guaranteed to be complete. This option makes invariant check-
    ing algorithm incomplete, therefore no "invariant is true" response can be
    given
-f Forces the building of the BDD FSM. Use this option if using option -d and
    verifying generalized invariants
-a Stop verification at the first property found false.
-u Change the semantics of the ";", operator from SEQUENCE to UNION.
-e "sere_expr" Provide the strategy from command line. This is an alternative to provide
    the strategy with an external file. In this case, the SERE formula must not
    begin with keyword 'grsequence'
-i sere_file Provide the strategy from file. This is an alternative to provide the strategy
    with the -e option. The SERE expression must start with the 'grsequence'
    keyword and must end with a ";"
```

```
msat_check_invar_bmc [-h | -n idx | -p "formula" | -P "name"] [-d
"mathsat" | "smtlib"] [-o filename] [-a alg] [-i] [-k max_len] [-e]
```

Performs invariant checking with BMC.
Command Options:

| -h | Shows a brief description of the available options |
| :---: | :---: |
| -u | Disables SMT solver invocation |
| -n idx | Checks the invariant (INVARSPEC) property specified with idx |
| -p "formula" | Checks the specified invariant property |
| -P name | Checks the invariant property with given name |
| -a alg | Uses the specified algorithm. Valid values are: <br> - classic <br> - een-sorensson <br> - falsification <br> - dual <br> - zigzag <br> - interp_seq <br> - interpolants |

-k max_len Maximum bound for BMC instead of using the variable bmc_length value. Use only when een-sorensson, falsification, dual or zigzag algorithm is selected
-i Use incremental version of falsification algorithm. Requires -a falsification
-e Performs an extra step for finding a proof. Can be used only with the eensorensson algorithm
-o filename Instead of checking the property the SMT problems are dumped into file filename with an additional suffix:
'_bmc_classic' for classic algorithm,
' bmc_base_n' for een-sorensson base problem
' bmc_step_n' for een-sorensson step problem where ' $n$ ' is the length of a path taken into account
-d format Selects the dumping format. Valid values are: "mathsat" and "smtlib"

Notice that if no property is specified, checks all LTL properties.

```
check_invar_bmc_itp [-h] | [-n idx | -p ``expr'' | -P ``name''] [-k
'`bound''] [-a '`alg'']
```

Performs invariant checking using interpolants based BMC algorithms. If no property is specified, checks all INVAR properties.
IMPORTANT: This command does not accept mixed integer and real types in the model's FSM constraints.

## Command Options:

| $-h$ | Shows a brief description of the available options. |
| :--- | :--- |
| -n number | Checks the property stored at the given index |
| -P name | Checks the property named name in the property database. |

```
-p "formula [IN cChecksthe"formula specified on the command-line. context is the module instance name which the variables in formula must be evaluated in.
-k idx Sets the BMC bound limit to be used.
-a alg Use the given algorithm for verification. Possible values are "mcmillan"
    (Default) and "itp_seq"
```

```
check_invar_ic3 [-h] [-d] [-i] [-O 0|1|2] [-a] [-g] [-x [-b]] [-m num]
[-v num] [-n number | -p "invar-expr" | -P "name"]
```

Checks invariant properties using the ic3 engines (simplic3 or smt)
For only the finite domain case it can check a property either internally or externally by invoking the simplic3 executable.

Environment variable "SIMPLIC3" is considered when searching for simplic3 external executable. If not specified, the executable is searched in the path specified when compiling the system.
When the domain is infinite (or when forced explicitly with option -i), msatic3 library is used to check the property.
IMPORTANT: This command does not accept mixed integer and real types in the model's FSM constraints.

Command Options:

```
-h
-d
-i
-O 0|1|2
```

Shows a brief description of the available options.
Disables the counterexample construction when proving false a property.
Forces the use of the engine for infinite domains (msatic3)
Only for finite: sets the preprocessing level (default: 2)
0 : No preprocessing.
1 : Enables sequencial preprocessing which searches equivalent latches and or constants
2 : Adds 2-step temporal decomposition to the preprocessor (default).

| $-a$ | Only for finite: enables abstract-refinement and BMC |
| :--- | :--- |
| -g | Only for finite: enables clause generalization, according to Zyad Has- <br> san, Aaron R. Bradley, Fabio Somenzi, "Better Generalization in IC3", <br> FMCAD'13 |
| -x | Invokes simplic3 as external executable instead of the internally linked li- <br> brary (works only for finite domains). |
| -b | When used with -x, dumps input file for simplic3 in binary format instead of <br> ASCII. |
| -v num | Enables verbosity. Default 0. Must be greater or equal to 0. <br> Only for finite: Max number of solvers to use for frames in IC3. Default 1. <br> Must be greater or equal to 1. |

```
-n number Checks the INVAR property with index number in the property database.
-p "invar-expr The command line specified invariant formula to be verified. context is
    [IN context]" the module instance name which the variables in invar-expr must be
                    evaluated in.
-P name Checks the INVAR property named name in the property database.
```

check_invar_local - Localized invariant checking
Command [F]

```
check_invar_local [-h] [-k] [-n idx | -p "expr" | -P "name"]
```

Performs invariant checking on the localized model. Model localization is performed before starting the checking process. This can greatly help in reducing resources (and time) required to check the property. Localization takes place using the property context as the input variable set.

Command Options:

$$
\begin{array}{ll}
-h & \text { Shows a brief description of the available options. } \\
-\mathrm{n} & \mathrm{idx} \\
-\mathrm{P} & \text { ''name'' } \\
-\mathrm{p} & \text { ''expr'' } \\
-\mathrm{k} & \\
& \begin{array}{l}
\text { Checks the property stored at the given index }
\end{array} \\
\text { Checks the formula specified on the command-line. }
\end{array}
$$

### 5.2.1 Incremental Cone Of Influence for Invariant Checking

In this section we list the commands for checking invariant properties that exploits abstraction based on incremental cone of influence reduction. The analysis starts considering the finite state transition corresponding to the cone of the property at distance 0 . If it succeeds in proving the property holds, it terminates. Otherwise, the counter-example is analyzed to see if it corresponds to a concrete counter-example. If the counter-example can be concretized, then we are done: the property is violated. Otherwise, the cone is refined adding new variables until either the property has been proved or disproved.
check_invar_inc_coi_bdd - BDD-based Incremental COI invariant checking

```
check_invar_inc_coi_bdd [-h] | [-n number | -p "invar-expr [IN context]"
| -P "name"] [-I]
```

Performs invariant checking using the BDD-based Incremental COI algorithm. Invariants to be verified can be provided as simple formulas (Only temporal operator allowed is "next") in the input file via the INVARSPEC keyword or directly at command line, using the option -p .

Option $-n$ or $-P$ can be used for checking a particular invariant of the model. If neither $-n$ nor -p nor -P are used, all the invariants are checked.

Command Options:

$$
\begin{aligned}
& \text {-p "invar-expr The command line specified invariant formula to be verified. context is } \\
& \text { [IN context]" the module instance name which the variables in invar-expr must be } \\
& \text { evaluated in. }
\end{aligned}
$$

$\underset{\substack{\text { Fondozione } \\ \text { BRUNO KESSER }}}{ }$

| $-n$ "idx" | Verifies the invariant with index "idx" within the Property Database |
| :--- | :--- |
| -P "name" | Verifies the invariant named "name" within the Property Database |
| -I | Execute traces over increasing size FSMs, based on Incremental COI |

check_invar_inc_coi_bmc - SAT-based Incremental COI invariant checking Command [F]

```
check_invar_inc_coi_bmc [-h] | [-n number | -p "invar-expr [IN context]"
| -P "name"] [-I] [-k bound]
```

Performs invariant checking using the SAT-based Incremental COI algorithm. Invariants to be verified can be provided as simple formulas (Only temporal operator allowed is "next") in the input file via the INVARSPEC keyword or directly at command line, using the option -p .
Option -n or -P can be used for checking a particular invariant of the model. If neither -n nor -p nor -P are used, all the invariants are checked.

Command Options:

$$
\begin{array}{ll}
\text {-p "invar-expr } & \text { The command line specified invariant formula to be verified. context is } \\
& {[\text { IN context] " the module instance name which the variables in invar-expr must be }} \\
\text { evaluated in. }
\end{array}
$$

```
msat_check_invar_inc_coi [-h] | [-n number | -p "invar-expr [IN
context]" | -P "name"] [-I] [-u] [-i] [-k bound]
```

Performs invariant checking using the SMT-based Incremental COI algorithm. Invariants to be verified can be provided as simple formulas (Only temporal operator allowed is "next") in the input file via the INVARSPEC keyword or directly at command line, using the option -p .

Option -n or -P can be used for checking a particular invariant of the model. If neither -n nor -p nor -P are used, all the invariants are checked.
IMPORTANT: In current implementation options -i and $-u$ are disabled, and an error is reported to the user when used.

IMIPORTANT: When using interpolation (option -i), integer and real types in the model's FSM constraints cannot be mixed.

Command Options:

```
-p "invar-expr The command line specified invariant formula to be verified. context is
    [IN context]" the module instance name which the variables in invar-expr must be
                    evaluated in.
-n idx Verifies the invariant with index "idx" within the Property Database
-P "name" Verifies the invariant named "name" within the Property Database
-I
    Execute traces over increasing size FSMs, based on Incremental COI
```

$\underset{\substack{\text { FoNDOZIONE } \\ \text { BRUNO KESSER }}}{ }$

| -u | Use unsat-cores variables for refinement (unsupported yet) |
| :--- | :--- |
| -i | Use interpolants variables for refinement (unsupported yet) |
| -k bound | The bound to be used for SMT algorithms |

check_invar_inc_coi - Incremental COI invariant checking
Command [F,I]

```
check_invar_inc_coi [-h] | -v eng -e eng [-n number | -p "invar-expr [IN
context]" | -P "name"] [-I] [-u] [-i] [-k bound]
```

Performs invariant checking using the Incremental COI algorithm. Invariants to be verified can be provided as simple formulas (Only temporal operator allowed is "next") in the input file via the INVARSPEC keyword or directly at command line, using the option -p .
Option $-n$ or $-P$ can be used for checking a particular invariant of the model. If neither $-n$ nor -p nor -P are used, all the invariants are checked.

IIMPORTANT: In current implementation options -i and $-u$ are disabled, and an error is reported to the user when used.

IIMPORTANT: For SMT, when using interpolation (option -i), integer and real types in the model's FSM constraints cannot be mixed.

Command Options:
-p "invar-expr
[IN context] " the module instance name which the variables in invar-expr must be
evaluated in.

### 5.3 Commands for LTL Model Checking

In this section we list the new commands for checking LTL properties. All these commands can be applied both to finite and infinite-state models (e.g. "ic3_check_ltlspec").

```
msat_check_ltlspec_bmc [-h | -n idx | -p "formula" | -P "name"] [-k
max_length] [-l loopback] [-d mathsat|smtlib] [-o filename]
```

Performs LTL checking with BMC.
Command Options:
\(\left.$$
\begin{array}{ll}\text {-h } & \begin{array}{l}\text { Shows a brief description of the available options } \\
\text {-u }\end{array} \\
\text {-t max_timespan } & \begin{array}{l}\text { Disables SMT solver invocation } \\
\text { Shows theory lemmas up to max_timespan } \\
-n ~ i d x ~\end{array}
$$ <br>

Checks the LTL property specified with idx\end{array}\right]\)| Checks the specified LTL property |
| :--- |
| -P name |
| -k max_length |
| -1 loopback | | Checks the LTL property with given name |
| :--- |
| Maximum bound for BMC instead of using the variable bmc_length value |
| Checks the property using loopback value instead of using the variable |

Notice that if no property is specified, checks all LTL properties.

## msat_check_ltlspec_sbmc_inc - LTL property check with Incremental SBMC

Command [I]

```
msat_check_ltlspec_sbmc_inc [-h | -n idx | -p "formula" | -P "name"] [-k
max_length] [-N] [-c]
```

Performs LTL incremental checking with SBMC.
Command Options:

| -h | Shows a brief description of the available options |
| :--- | :--- |
| -n idx | Checks the LTL property specified with idx |
| -p "formula" | Checks the specified LTL property |
| -P name | Checks the LTL property with given name |
| -k max_length | Maximum bound for BMC instead of using the variable bmc_length value <br> -N |
| -C | Does not perform virtual unrolling |
| Performs completeness check |  |

Notice that if no property is specified, checks all LTL properties.

```
check_ltlspec_klive [-h] [-d] [-m num] [-i] [-x] [-g] [-b] [-0 [0|1]]
[-e] [-E num] [-L] [-v num] [-n number | -p "ltl-expr" | -P "name"]
```

Checks LTL properties using the ic3 engines (simplic3 or smt) using K-Liveness.
For only the finite domain case it can check a property either internally or externally by invoking the simplic3 executable.

Environment variable "SIMPLIC3" is considered when searching for simplic3 external executable. If not specified, the executable is searched in the path specified when compiling the system.

When the domain is infinite (or when forced explicitly with option -i), msatic3 library is used to check the property.
IIMPORTANT: This command does not accept mixed integer and real types in the model's FSM constraints.
$\underset{\substack{\text { Fovedzione } \\ \text { BRINO KESSER }}}{ }$

## Command Options:

| -h | Shows a brief description of the available options. |
| :---: | :---: |
| -d | Disables the counterexample construction when proving false a property. |
| -i | Forces the use of the engine for infinite domains (msatic3) |
| -0 011 | Only for finite: sets the preprocessing level (default: 1) |
|  | 0 : No preprocessing. |
|  | 1 : Enables sequencial preprocessing which searches equivalent latches and or constants (default). |
| -g | Only for finite: enables clause generalization, according to Zyad Hassan, Aaron R. Bradley, Fabio Somenzi, "Better Generalization in IC3", FMCAD' 13 |
| $-x$ | Invokes simplic3 as external executable instead of the internally linked library (works only for finite domains). |
| -b | When used with -x, dumps input file for simplic3 in binary format instead of ASCII. |
| -e | Only for finite: enables extraction additional liveness stabilizing constraints in preprocessing generalization. |
| -E num | Only for finite: number of candidates to consider for liveness constraint extraction.(0: disabled, 3000 default). |
| -L | Only for finite: disables complementation of k-liveness with BMC (if disabled no counterexample can be computed). |
| -m | Only for finite: max number of solvers to use for frames in IC3. Default 1. Must be greater or equal to 1 . |
| -v <num> | Enables verbosity. Default 0 . Must be greater or equal to 0 . |
| -n number | Checks the LTL property with index number in the property database. |
| -p "invar-expr <br> [IN context | The command line specified LTL formula to be verified. context is the module instance name which the variables in invar-expr must be evaluated in. |
| -P name | Checks the LTL property named name in the property database. |

```
check_ltlspec_simplify [-h] [-n index | -p '`prop'' | -P '`name''] [-s]*
[-r]*
```

Performs model checking of LTL formulas. LTL model checking is reduced to CTL model checking as described in the paper by [CGH97a].

The model on which the model checking is performed is simplified using the Model Simplifier and the Range Reduction systems.
By default, Model Simplification and Range Reduction are used, but a chain of simplifications to be performed over the model can be specified using the $-s$ and the $-r$ command options.
A ltl-expr to be checked can be specified at command line using option - p . Alternatively, options -n and $-P$ can be used for checking a particular formula in the property database. If neither $-n$ nor -p nor -P are used, all the LTLSPEC formulas in the database are checked.

## Command Options:

```
-p '`prop''
-P "name"
-n index
-s
-r
```

An LTL formula to be checked.
Checks the LTL property named "name"
Checks the LTL property with index index in the property database.
Adds Model Simplification to the chain of simplifications. This option can be used multiple times
-r Adds Range Reduction to the chain of simplifications. This option can be used multiple times

### 5.3.1 Incremental Cone Of Influence for LTL Model Checking

In this section we list the commands for checking LTL properties that exploits abstraction based on incremental cone of influence reduction. Similarly to the case of verification of invariants, the analysis starts considering the finite state transition corresponding to the cone of the property at distance 0 . If it succeeds in proving the property holds, it terminates. Otherwise, the counter-example is analyzed to see if it corresponds to a concrete counterexample. If the counter-example can be concretized, then we are done: the property is violated. Otherwise, the cone is refined adding new variables until either the property has been proved or disproved.

```
checkltlspec_inc_coi_bdd - BDD-based Incremental COI LTL properties
```

check_ltlspec_inc_coi_bdd [-h] | [-n number | -p "ltl-expr [IN context]"
| -P "name"] [-I]

```

Performs LTL checking using the BDD-based Incremental COI algorithm. LTL properties to be verified can be provided as LTL formulas in the input file via the LTLSPEC keyword or directly at command line, using the option -p .

Option -n or -P can be used for checking a particular LTL property of the model. If neither -n nor -p nor \(-P\) are used, all the LTL properties are checked.

Command Options:
\[
\begin{array}{ll}
\text {-p "ltl-expr } & \text { The command line specified LTL formula to be verified. context is the } \\
& \text { [IN context] " module instance name which the variables in ltl-expr must be evaluated } \\
\text { in. }
\end{array}
\]
```

check_ltlspec_inc_coi_bmc - SAT-based Incremental COI LTL properties check-

```
check_ltlspec_inc_coi_bmc [-h] | [-n number | -p "ltl-expr [IN context]"
| -P "name"] [-I] [-k bound]
```

Performs LTL checking using the SAT-based Incremental COI algorithm. LTL properties to be verified can be provided as LTL formulas in the input file via the LTLSPEC keyword or directly at command line, using the option -p .

Option -n or -P can be used for checking a particular LTL property of the model. If neither -n nor -p nor $-P$ are used, all the LTL properties are checked.

Command Options:

$$
\begin{array}{ll}
\text {-p "ltl-expr } & \text { The command line specified LTL formula to be verified. context is the } \\
& \\
& \\
-\mathrm{n} \text { IN context] " module instance name which the variables in ltl-expr must be evaluated } \\
\text { in. } & \text { Verifies the LTL property with index number within the Property Database } \\
-\mathrm{P} & \text { "name" } \\
-\mathrm{I} & \\
\text { Verifies the LTL property named "name" within the Property Database }
\end{array}
$$

msat_checkltlspecinc_coi - SMT-based Incremental COI LTL properties
Command [I] checking

```
msat_check_ltlspec_inc_coi [-h] | [-n number | -p "ltl-expr [IN
context]" | -P "name"] [-I] [-u] [-i] [-k bound]
```

Performs LTL checking using the SMT-based Incremental COI algorithm. LTL properties to be verified can be provided as LTL formulas in the input file via the LTLSPEC keyword or directly at command line, using the option -p .
Option -n or -P can be used for checking a particular LTL property of the model. If neither -n nor -p nor -P are used, all the LTL properties are checked.
IMPORTANT: In current implementation options - i and - $u$ are disabled, and an error is reported to the user when used.

IMPORTANT: When using interpolation (option -i), integer and real types in the model's FSM constraints cannot be mixed.

Command Options:

```
-p "ltl-expr The command line specified LTL formula to be verified. context is the
    [IN context]" module instance name which the variables in ltl-expr must be evaluated
                    in.
-n idx Verifies the LTL property with index "idx" within the Property Database
-P "name" Verifies the LTL property named "name" within the Property Database
-I Execute traces over increasing size FSMs, based on Incremental COI
-u Use unsat-cores variables for refinement
-i Use interpolants variables for refinement
-k bound The bound to be used for SMT algorithms
```

check_ltlspec_inc_coi - Incremental COI LTL properties checking

```
check_ltlspec_inc_coi [-h] | -v eng -e eng [-n number | -p "ltl-expr [IN
context]" | -P "name"] [-I] [-u] [-i] [-k bound]
```

Performs LTL checking using the Incremental COI algorithm. LTL properties to be verified can be provided as LTL formulas in the input file via the LTLSPEC keyword or directly at command line, using the option -p . Option -n or -P can be used for checking a particular LTL property of the model. If neither -n nor -p nor -P are used, all the LTL properties are checked.
IMPORTANT: In current implementation options $-i$ and $-u$ are disabled, and an error is reported to the user when used.

IIMPORTANT: For SMT, when using interpolation (option -i), integer and real types in the model's FSM constraints cannot be mixed.

Command Options:

$$
\begin{aligned}
& \text {-p "ltl-expr The command line specified LTL formula to be verified. context is the } \\
& \text { [IN context] " module instance name which the variables in ltl-expr must be evaluated } \\
& \text { in. }
\end{aligned}
$$

### 5.3.2 Compositional Reasoning for LTL Model Checking

Compositional model checking is a verification method that aims at reducing the verification problem of large systems to smaller, possibly localized verification problems. This is done in order to try to avoid the "state explosion problem". When reasoning compositionally about two systems $A$ and $B$, it is often necessary to assume the correctness of $A$ to verify $B$, and vice-versa. In the literature this "apparent" circularity has been resolved by induction over time. The induction over time is made explicit by assuming that a property $P$ only up to time $t-1$ when proving $Q$ at time $t$, and vice-versa. The proof obligations incurred using this method can be discharge with model checking. This approach has been described in [McM99]. In this section we describe the commands for performing the above outlined compositional reasoning.

```
checkltlspec_compositional - Circular compositional reasoning model check-
    Command [F,I]
ing
```

```
check_ltlspec_compositional [-h] | -f ''proof-file'' [-n '`node''] [-t
    ''check-technique'' ]
```

Performs circular compositional reasoning model checking as described in [McM99] and [PK99].
An assertion is a condition that must hold true in every possible execution of the program. Assertions refer to properties in LTL.

In order to apply the circular compositional rule, one has to supply the set of assertions to be proved and the proof graph. From these, a sufficient set of proof obligations on the form of LTL formulas are built.
A property is specified using the following syntax:
name : assert $\langle$ formula $\rangle$;
When specifying the proof graph, properties are refererred to by their names. An arc (p1, p2) in the proof graph is specified as follows:
using n 1 prove n 2 ;
where n 1 and n 2 are the respective names of properties p 1 and p 2 . A list of assumption can also be used when verifying a property, specifying a comma-separated list of assumptions.
using n1, $\mathrm{n} 2, \mathrm{n} 3$ prove n 4 ;
Such a "proof" may not contain circular chains of reasoning. The system verifies that every cycle in the proof graph is cut by a unit delay arc. A unit delay arc is specified by putting the assumption in parentheses, as follows:
using ( n 1 ) prove n 2 ;
$\underset{\substack{\text { Fondazione } \\ \text { BRINO } \\ \text { KESS ER }}}{ }$

Command Options:

| -t technique | Use the specified technique to perform model checking. Valid techniques <br> are $\{b d d, b m c, s m t\}$ |
| :--- | :--- |
| -n node | Perform the check only for the specified node instead of checking all the <br> nodes |
| $-£$ proof-file | Reads the proof graph from the specified file |

### 5.4 Commands for Computing Reachable States

## compute_reachable_guided - Guided reachability set of reachable states

```
compute_reachable_guided [-s] [-S] [-R] [-P] [-d] [-u] (-h | -e
"sere_expr" | -i sere_file)
```

Computes the set of reachable states of the given model using Guided Reachability algorithm over the given strategy. By default the computation is done in two steps. At first, states satisfying the strategy are computed until fixpoint is reached. Then the forward image is applied to the computed states independently of the strategy until all the reachable states are detected.
If the set of reachable states has already been computed, the command returns immediately since there is nothing more to compute.
The resulting reachable states are used to simplify image and preimage computations in consecutive model checking commands. This can result in improved performance on models with sparse state spaces.
The strategy must be a valid PSL formula. Allowed PSL operators are: "; $[*][* N]($ with $N>0) \mid "$

## Command Options:

\(\left.\begin{array}{ll}-s \& Uses model simplification over the given model <br>
Enables the use of a further simplified FSM for each atomic part of the given <br>

SERE\end{array}\right]\)| Enables the use of Implicit Frame Condition for each atomic part of the given |
| :--- |
| SERE |

### 5.5 Commands for Reasoning via Abstraction

Predicate abstraction is a technique that is used to prove properties of finite- and infinite-state systems. It is a combination of theorem proving and model checking techniques. Given a concrete finite- or infinite-state system and a set of predicates, a conservative finite state abstraction is generated. (For every execution in the concrete system there is a corresponding execution in the abstract system.) The abstract version of the verification condition is model checked in this abstract system. If the property is verified then it holds in the concrete system. Otherwise an abstract counter-example trace is generated. There could be a concrete counter-example corresponding to it, in which case there is a bug in the design, or the abstract counter-example is an artifact of the abstraction. The counter-example can be analyzed to find a real bug or to suggest extra predicates to refine the abstraction thereby avoiding that particular spurious trace. Then the process starts anew. The abstraction refinement process is guaranteed to terminate for finite-state systems (if resources permits). However, the process is not guaranteed to terminate for infinite-state systems: proving arbitrary (safety) properties of an infinite state system is not decidable, but for a large number of problems this method can successfully prove properties.

In NUXMV we provide two complementary approaches, both based on predicate abstraction. The first relies on the explicit computation of the abstract transition system. The second, uses an implicit abstraction to avoid the expensive computation of the abstract transition system.

### 5.5.1 Explicit Predicate Abstraction

These commands implements the functionalities for performed Counterexample Guided predicate Abstraction Refinement (CEGAR) [CGJ ${ }^{+}$03]. The CEGAR approach requires the computation of a quantifier-free formula that is equivalent to the abstract transition relation w.r.t. a given set of predicates. This, in turn, requires the solving of an ALLSAT problem [LNO06]. For this step, NUXMV implements different techniques: the combination of BDD and SMT $\left[\mathrm{CCF}^{+} 07, \mathrm{CFG}^{+} 10\right]$, where BDDs are used as compact Boolean model enumerator within an AllSMT approach; the technique that exploits the structure of the system under verification, to partition the abstraction problem into the combination of several smaller abstraction problems [CDJR09]. For the refinement step to discard the spurious counterexample, nUXMV implements three approaches based on the analysis of the unsatisfiable core, on the analysis of the interpolants, and on the weakest preconditions..

The command "config_abstraction" sets the options that control how the abstraction is performed. The command "add_abstraction_preds" allows to specify the predicates to be used for CEGAR. The command "build_abstract_model" computes the abstract transition system w.r.t. the specified set of predicates. Then, the model can be dumped into a file with the command "write_abstract_model". The command "check_invar_cegar_predabs" performs the CEGAR loop checking the given property.
config_abstraction - Configures the options for abstraction computation

```
config_abstraction [-h] [-e <output>] [-d <output>] [-a <engine>] [-c
(0|1)] [-t (0|<number>)] [-b (0|1)] [-s]
```

This command sets the options for abstraction computation using the build_abstract_model command.

Command Options:

$$
\begin{array}{ll}
\text {-h } & \text { Shows the online help message. } \\
\text {-a engine } & \text { The abstraction engine to be used. The parameter engine can be } \\
\text { msat (AllSMT) or bdd }\left[C F G^{+} 10\right] \text { or bddarray }\left[\mathrm{CFG}^{+} 10\right] \text { or bool or } \\
\text { structural [CDJR09]. }
\end{array}
$$

$\underset{\substack{\text { Fondazione } \\ \text { BRINO } \\ \text { KESS ER }}}{ }$

add_abstraction_preds - Extracts the abstraction precision from a model or
Command [I]
from an external file

```
add_abstraction_preds [-h] [-a | -p number | -m number | -i file] [-o
file] [-s]
```

Extracts and/or shows the abstraction precision. The precision is the set of mirror variables and predicates to be used in the abstraction. The precision can be specified in two ways. First, mirrors and predicates can be declared in the model using the MIRROR and PRED keywords. A second possibility is to specify the list of predicates and mirrors in a separate file and use the -i option. The file format is the following. The file starts with the string PREDICATES followed by an arbitrary list of either PRED [predicate] or MIRROR [mirror variable]. The extracted precision is then used in the commands build_abstract_model and check_invar_cegar_predabs.

Command Options:

| -h | Shows the online help message. |
| :--- | :--- |
| -a | Adds all predicates and mirrors from model. |
| -p number | Adds the predicate with the specified number from model. |
| $-m$ number | Adds the mirror with the specified number from model. |
| $-i$ file | Reads the precision specified in the given file. |
| $-o$ file | Writes the precision extracted so far in the specified file. |
| $-s$ | Shows the precision extracted so far. |

```
build_abstract_model [-h]
```

This command computes and internally stores the abstraction of the model given the precision extracted using add_abstraction_preds with the options set by config_abstraction. The generated FSM can be dumped using write_abstract_model and disposed using quit_abstraction.

Command Options:
-h Shows the online help message.

```
quit_abstraction [-h]
```

This command disposes the FSM computed by build_abstract_model in order to allow for another one to be generated (with different options or with a different precision).


Command Options:
-h Shows the online help message.
write_abstract_model - Dumps the abstracted FSM to a file
Command [I]
write_abstract model [-h] <filename>
This command dumps the FSM computed by build_abstract_model to the given filename.
Command Options:
-h
Shows the online help message.

## check_invar_cegar_predabs - Checks a property using CEGAR

Command [I]

```
check_invar_cegar_predabs [-h] [-n <num> | -P <name> | -p <formula>] [-l
<num>]
```

This command sets performs the model checking of a property using the CEGAR loop. The options for the abstraction phase can be set using config_abstraction command. The property to be checked can be specified by its number ( -n parameter), by its name ( -P parameter) or by entering an invariant formula ( -p option). The number of abstraction-refinement steps is bounded by the -1 option and by default it is unlimited It is possible to specify predicates and mirrors to be used in all the abstractions cycles by using the add_abstraction_preds command.

Command Options:

$$
\begin{array}{ll}
-\mathrm{h} & \text { Shows the online help message. } \\
-\mathrm{n} \text { num } & \text { The number of the property to be checked. } \\
-\mathrm{P} \text { name } & \text { The name of the property to be checked. } \\
-\mathrm{p} \text { formula } & \text { Adds an invariant property with the given formula and checks it. } \\
-1 \text { num } & \text { Bounds the number of cycles to num. }
\end{array}
$$

### 5.5.2 Implicit Predicate Abstraction

We also complement the CEGAR based predicate abstraction algorithms with new algorithms that combine abstraction with BMC and k-induction [Ton09]. The algorithms do not rely on quantifier elimination techniques to compute the abstraction, but encode the model checking problem over the abstract state space into SMT problems. The advantage, is that they avoid the possible bottleneck of abstraction computation.

```
msat_check_invar_bmc_implabs - Verifies invariant properties using BMC in
```

msat_check_invar_bmc_implabs [-h] [-k bound] [-a] (-n prop_index | -P
prop_name | -p formula)

```

Checks invariant properties using bounded model checking with k -induction in combination with abstraction w.r.t. a given set of predicates specified in the input file. The technique that combines abstraction and k -induction has been described in [Ton09].
\(\underset{\substack{\text { Fondozione } \\ \text { BRUNO KESSER }}}{ }\)

Command Options:
```

-h
-k bound
-n num
-P prop_name
-p formula
-a

```

Shows a brief description of the available options.
Specifies the maximum bound for bounded model checking and k-induction. Default value is the one specified by variable bmc_length. If value 0 is used, the algorithms continues to increment k until either the property has been proved, or a counterexample has been found, or the resources are exhausted.

Specifies the id of an invariant property. If the id does not corresponds to an invariant property, then an error is issued.
Specifies the name of an invariant property. If the name does not corresponds to an invariant property, then an error is issued.
Specifies an invariant property. It will be added to the property database.
Disables the use of abstraction, and simply performs the verification using the standard bounded model checking algorithms. The specification of one invariant property is mandatory.
```

msat_check_invar_bmc_cegar_implabs [-h] | [-k bound] [-r meth] [-i meth]
[-s] [-m] [-d] [-c] [[-n prop] | [-p "invar"] | [-P name]]

```

Performs invariant checking with implicit abstract model checking as described in [Ton09]. This technique does not compute the abstraction explicitly like it is the case in classical Counterexample Guided Abstraction Refinement (CEGAR). This would save resources (memory and time) to check the property. If neither of -o, -p , or -P is specified, the command tries to

Command Options:
```

-k bound

```
-r meth
-i meth

Shows a brief description of the available options.
Specifies the maximum bound for bounded model checking and k-induction. Default value is the one specified by variable bmc_length. If value 0 is used, the algorithms continues to increment k until either the property has been proved, or a counterexample has been found, or the resources are exhausted.
\(-r\) meth Refinement method for generating new predicates. (a:automatic (default), m:manual, h: hybrid). automatic : Ignore predicates in smv file and generate new predicates manual : Take predicates in smv file and do not generate any new predicates hybrid : Take predicates in smv file and generate new predicates
K -induction method for termination condition. (f:full(BW+FW) (default), b:backward only, n:none).
full : Check backward induction first and then forward induction; backward : Only check backward condition; none : None. This is fully Abstract BMC. Program will only terminate if it reaches the bound k or find a bug.
\begin{tabular}{ll}
\(-s\) & Incrementally add simulation condition. \\
-m & Tries to minimize the number of predicates added during the search. \\
-d & Enables for the discovery of invariants during the search. \\
-c & \begin{tabular}{l} 
Enables for the fresh restart after each refinement step.
\end{tabular} \\
-n prop & \begin{tabular}{l} 
Checks the property stored at the given index, assuming it is an invariant. If \\
it is not an invariant, an error is issued.
\end{tabular} \\
-p & 'invar''
\end{tabular} \begin{tabular}{ll} 
The command line specified invariant formula to be verified. context is
\end{tabular}

\subsection*{5.6 Commands for Format Conversions}

This subsection contains commands for converting the nUXMV format into other external formats.

\subsection*{5.6.1 Commands for aiger 1.9.4 format support}
aiger 1.9.4 is a format, library and set of utilities for And-Inverter Graphs (AIGs) [BHW11]. The aiger 1.9.4 format has an ASCII and a binary version. As described in the documentation of aiger 1.9.4 [BHW11], the ASCII version is the format of choice if an AIG is to be saved by an application which does not want to use the aiger 1.9.4 library. We refer the reader to the aiger 1.9.4 [BHW11] documentation for details about the format.

In this section we describe the commands for reading the aiger 1.9.4 format (both in the ASCII and in the binary formats). Moreover, we also describe the command to dump a NUXMV model without Reals and Integers into a set of aiger 1.9.4 files (one for each property).
read_aiger_model - Reads and loads an aiger model
```

read_aiger_model [-h] | [-i filename] [-r] [-m]

```

The command imports a model in aiger 1.9.4 format into nUXмV. The loaded model can then be used as any other input file in the extended language accepted by the tool.
If the aiger file is in the aiger 1.9.4 format with both bad, justice, fairness do not consider the output as properties. Each fairness conditions \(f_{i}\) is transformed in FAIRNESS \(f_{i}\). Each bad conditions \(b_{i}\) is transformed in INVARSPEC \(!b_{i}\). Each justice constraint \(J_{i}=j_{0}^{i}, \ldots, j_{n}^{i}\) is transformed in LTLSPEC ! \(\bigwedge_{j_{k}^{i} \in J_{i}} G F_{k}^{i}\).
On the other hand, if the input file does not contain neither fairness, nor justice and nor bad, than if there is a single output \(o_{0}\) it is interpreted as an INVARSPEC \(!o_{0}\). Otherwise, if more than one output is specified, i.e. \(O=o_{0}, \ldots, o_{n}\), each output \(o_{i}\) is interpreted as a \(G F o_{i}\) and the corresponding property is LTLSPEC \(!\bigwedge_{o_{i} \in O} G F o_{i}\).

Command Options:
\[
\begin{aligned}
& -h \\
& -i \text { filename } \\
& -r \\
& -m
\end{aligned}
\]

Shows a brief description of the available options.
Reads and loads the model from "input-file".
Builds a relational hierarchy instead of building a functional one, i.e. uses INIT / TRANS instead of ASSIGN
-m Uses monitor variables recognition, which tries to detect which variables in the aiger model are monitor/support variables, and on success, slightly reduces the number of variables and simplifies the Transition Relation. This should work almost every time when loading "write_aiger_model"-generated files.
write_aiger_model - Dump of the current model in aiger format
```

write_aiger_model [-h] | -p "prefix" | -f "output" [-i | -l] [-n index]
[-b] [-d path]

```

Dumps the currently loaded model in aiger format using the aiger 1.9.4 format. Input format before aiger 1.9.4 is only supported for reading (see read_aiger_model for details).

If \(-p\) "prefix" is specified, a various number of aiger 1.9.4 files is generated, one for each property of kind LTL or INVARSPEC in the property database. (Other types of properties are not supported.) Each file will be named "prefix_proptype_propidx.[aig|aag]". Each generated file represents a model checking instance for the corresponding property.

If option -f "output" is specified instead, one file named "output" will be generated, which represents the Finite State Machine of the input model only without properties.

Command Options:
```

-h Shows a brief description of the available options.
-b Dumps the output files in binary format instead of ASCII format
-i Dumps models only for invariant properties
-1 Dumps models only for LTL properties.
-n index Dumps models only for the property at index n.
-d path The directory where to save files. Default is "."
-p prefix Dumps one model foreach property. Each generated file will be named "pre-
fix_proptype_propidx.[aig|aag]"

```

\subsection*{5.6.2 Commands for VMT format support}

The VMT format is an extension of the SMT-LIBv2 [BST12] (SMT2 for short) format to represent symbolic transition systems.

VMT exploits the capability offered by the SMT2 language of attaching annotations to terms and formulas in order to specify the components of the transition system and the properties to verify. More specifically, the following annotations are used:
: next name is used to represent state variables. For each variable \(x\) in the model, the VMT file contains a pair of variables, \(x^{c}\) and \(x^{n}\), representing respectively the current and next version of \(x\). The two variables are linked by annotating \(x^{c}\) with the attribute : next \(x^{n}\). All the variables that are not in relation with another by means of a : next attribute are considered inputs.
:init true is used to specify the formula for the initial states of the model. This formula should contain neither next-state variables nor input variables. (The "dummy" value true in the annotation is needed because the current SMT2 standard requires annotations to always have an associated value.)
:trans true is used to specify the formula for the transition relation.
: invar-property \(\boldsymbol{i d x}\) is used to specify invariant properties, i.e. formulas of the form \(G p\), where \(p\) is the formula annotated with : invar-property. The non-negative integer idx is a unique identifier for the property.
: live-property \(i d x\) is used to specify an LTL property of the form \(F G p\), where \(p\) is the formula annotated with : live-property. The non-negative integer \(i d x\) is a unique identifier for the property.

In a VMT file, only annotated terms and their sub-terms are meaningful. Any other term is ignored. Moreover, only the following commands are allowed to occur in VMT files: set-logic, set-option, declare-sort, define-sort, declare-fun, define-fun.(For convenience, an additional (assert true) command is allowed to appear at the end of the file.)
The following example shows a simple nUXmV model (left) and its corresponding VMT translation (right).
\begin{tabular}{|c|c|}
\hline NUXMV & VMT \\
\hline \begin{tabular}{l}
-- this is a comment MODULE main \\
VAR x : integer; \\
INIT \(\mathrm{x}=1\); \\
TRANS next \((\mathrm{x})=\mathrm{x}+1\); \\
INVARSPEC x > 0;
\end{tabular} & ```
; this is a comment
(declare-fun x () Int)
(declare-fun xn () Int)
(define-fun .sv0 () Int (! x :next xn))
(define-fun .init () Bool (! (= x 1) :init true))
(define-fun .trans () Bool (! (= xn (+ x 1)) :trans true))
(define-fun .p0 () Bool (! (> x 0) :invar-property 0))
``` \\
\hline
\end{tabular}

Since the SMT2 format (and thus also the VMT one that inherits from SMT2) does not allow to annnotate the declaration of variables, it is a good practice to insert immediately after the declaration of the variables a set of defines to specify the relations among variables. See for instance the define .sv0 in the example above that introduces the relation between x and \(\mathrm{x} n\).
In the distribution of the NUXMV (within directory contrib), we also provide conversion scripts from other formats (e.g. the from the BTOR language of Boolector [Boo] to the language of the NUXMV and vice-versa.
write_vmt_model - Dumps the model with a single property in VMT format
Command [I]
```

write_vmt_model [-h] [-o filename] [-n prop_number | -i "invar_expr" |
-l "ltl_expr"]

```

Dumps the model with the specified property in VMT format. VMT is an extension of the SMT-LIBv2 format for specifying symbolic transition systems.

Command Options:
```

-h Shows a brief description of the available options.
-o filename The filename where to dump the model and the possible LTL or invariant
property.
-n prop_number Index of the property to dump in the property database. Only invariants and
LTL properties are supported.
-i "invar_expr" An expression specifying the invariant property to dump. The property is
also added to the property database.
-l "ltl_expr" An expression specifying the LTL property to dump. The property is also
added to the property database.

```

\subsection*{5.7 Commands for Model Transformation}

In this section we report a set of commands that could be used to generate simplified models, and to explore the model e.g. generating XMI format.

\subsection*{5.7.1 Commands for Model Simplification}
```

write_simplified_model_rel [-h] [-o filename] [-e '`expr'']* [-l
''prop'']* [-i ''prop'']* [-c ''prop'']* [-I | -D] [-r]

```

Writes the currently loaded SMV model in the specified file, after having flattened and simplified it. INVARs and ASSIGNs are taken as assumptions for simplification. Those expressions are processed in order to find strong dependencies between variables, so that some of them can be simply removed, reducing the space search. For example, having "INVAR \(a=b\) " means that all occurrences in the input model of one of the two variables involved in the expression can be replaced with the other one. Also inlining is applied as much as possible in order to reduce the expressions size.
During simplification, processes are eliminated and equivalent structures are set up.
If no file is specified, resulting simplified flat model is dumped to standard output.
Command Options:
\begin{tabular}{ll}
-h & Shows a brief description of the available options. \\
-o filename & Attempts to write the simplified SMV model in filename \\
-e expr & Adds the given assumption to the simplifier \\
-l prop & Adds the given LTL property to the simplifier \\
-i prop & Adds the given INVAR property to the simplifier \\
-c prop & Adds the given CTL property to the simplifier \\
-D & Does not add any define declaration to the output model \\
-I & Disable defines and array defines inlining \\
-r & Disable properties rewriting
\end{tabular}
```

write_simplified_model_func [-h] | [-o filename] [-d] [-D]

```

Writes the currently loaded SMV model in the specified file, after having flattened and simplified it. Assignments of the form \(\mathrm{A}:=\mathrm{B}\), in which A is a variable and B is a constant or a variable are transformed into defines, thus reducing the state space of the model. Additionally, when 2 variables are assigned the same expression then one of variables is converted to a define equal to the second variable. Assignments to init, current and next variables are taken into account. Daggification can be used in order to share common subformulas.
If no file is specified, resulting reduced simplified model is dumped to standard output.
Command Options:
\[
\begin{array}{ll}
\text {-o filename } & \begin{array}{l}
\text { Attempts to write the simplified SMV model in filename } \\
\text {-D } \\
\text { Enables daggification and simplification of expressions to make the detection } \\
\text { of equivalent variables more effective }
\end{array} \\
-\mathrm{d} & \begin{array}{l}
\text { Disables optimization that creates one fresh variable to distinguish the first } \\
\text { state for assignments }
\end{array}
\end{array}
\]
```

write_range_reduced_model [-h] | [-c] [-d] [-g] [-o file] [-f fixp]

```

Writes the currently loaded SMV model in the specified file, after having flattened it and reduced its variable ranges. Processes are eliminated and a corresponding reduced model is printed out.
If no file is specified, resulting reduced flat model is dumped to standard output.

\section*{Command Options:}
```

-h

```
-c Enables detection of counters from data clusters.
-d Disables normal range detection (counters still may be detected).
-g Adds invariant about guessed ranges to the property database.
-o file Attempts to write the flat SMV model in file
-f fixp Sets the fixpoint to be used for range extraction. Default is 20. Must be a
non-negative integer
```

build_simplified_property [-h] | [-a] | [ [-n <index>] | [-N <name>] [-c
<ctlspec>] | [-l <ltlspec>] | [-i <invarspec] ]* [-e]

```

Performs property simplifications on a set of properties. INVARs and ASSIGN are taken as assumptions for simplification, as done for command write_simplified_model_rel. During simplification, processes are eliminated and equivalent structures are set up. Each simplified property is associated to a simplified model and registered in the property database, so that future calls to usual Model Checking commands (e.g. check_invar) will verify freshly created properties using the simplified model.
Using build_simplified_property with input model \(M\) and then performing model checkin, is actually equivalent to reading model \(M\), dumping the simplified version of \(M, M^{\prime}\), using command write_simplified_model_rel, reading \(M^{\prime}\) and finally perform model checking.

Command Options:
```

-h Shows a brief description of the available options.
-a Selects all registered properties for simplification.
-l prop Specifies a LTLSPEC property to simplify.
-i prop Specifies an INVARSPEC property to simplify.
-c prop Specifies a CTLSPEC property to simplify.
-e expr Assume given expression.
-n number Simplifies property with index number in the property database.
-N name Simplifies property with name name in the property database.

```
write_hier_coi_model - Localize a model
Command [I]
write_hier_coi_model [-h] | -i iv-file [-k] [-o filename]
Given an input variables description file, this commands creates a localized version of the current model, that is restricted to the parts of the model which depend on the variables given as input. The optional parameter keep behavior forces adding the dependencies of the constraints in which a variable in input set occurs.

Command Options:
\[
\begin{aligned}
& \text {-h } \\
& \text {-i iv_file }
\end{aligned}
\]

Shows a brief description of the available options.
Reads the variable names from the specified "iv_file" instead of searching in the model matching the counter structure. Input file may contain variable names and/or instances with the intended meaning to include all variables within the instance
\begin{tabular}{ll}
-k & \begin{tabular}{l} 
This flag enables recursive dependencies resolving when performing simpli- \\
fication, thus resulting in a stricter, behavior-preserving, approximation.
\end{tabular} \\
\(-\bigcirc\) filename & Writes the output generated by the command in to the file filename.
\end{tabular}
write_countacc_model - Create "accelerated" models.
Command [I]
```

write_countacc_model [-h] | [-c] [-o filename] [-i iv_file] [-l
list_file] [-v] [-V] [-s] [-p]

```

This commands creates an "accelerated" version of the current model changing the behavior of its counter variables. A counter is a word variable that is initialized to 0 , can be enabled by an arbitrary boolean expression and that when enabled, increases by 1 at each step until a limit value is reached or the enabling condition is false. When a counter reaches the limit or is disabled it is reset to 0 . The accelerated model is such that in a single step counters possibly increase their value by a number greater than 1 .

Note that the accelerated model does not preserve all properties of the model except invariants.

\section*{Command Options:}
\(\left.\begin{array}{ll}\text {-h } & \text { Shows a brief description of the available options. } \\ -\mathrm{c} & \text { Performs a check on the constraints on counter variables. } \\ -0 \text { filename } & \begin{array}{l}\text { Writes output to "filename" instead of stdout. } \\ \text { Read the counter names from the specified "iv_file" instead of searching in } \\ \text { the model matching the counter structure. }\end{array} \\ \text { This option is incompatible with the option -l. } \\ \text { Read the counter names and limit values from the specified "list_file" instead } \\ \text { of searching in the model matching the counter structure. The file must be } \\ \text { in the following format: }\end{array}\right]\)

\subsection*{5.7.2 Commands for Model Exploration}
write_xmi_model [-F fsm] [-o filename] [-a explist] [-f format]
Converts the symbolic FSM representing the model to an explicit finite state machine (EFSM). Then prints it in XMI format to the file specified with the \(-\circ\) option.
If the option -a is not used, all finite variables of the model are taken into consideration.
Otherwise, the EFSM is abtstracted to the specified expressions: the states will be made of the specified expressions only. The expressions can be boolean, or single variables with boolean, integer, enumeration or word type. Expressions made by a single variable are expanded to all the valid assignments of the variable. This behavior may be critical with words variables, that could easily have a wide range.

Command Options:
```

-h
-F fsm

```
-a expr_list Abstract the EFSM to the specified expressions. Explist is "exp_1, ..., exp_n"
where every exp_i is a variable or a boolean expression.
-o file Redirects the output to the specified file; default: standard output.
\(-f\) format Format the XMI in a specific way. Currently, the only valid value is "ea", for
making the xmi readable by Enterprise Architect.

\subsection*{5.8 NUXMV environment variables}

In this section we describe all the environment variables that may affect the behavior of the new features of nuXmv.
```

abstraction.engine
Environment Variable

```

Specifies the engine to be used while computing the abstraction during the CEGAR loop. Possible values for this variable are:
- msat: (default) This approach uses AllSMT [LNO06] to compute the abstraction. It assumes all the predicates and mirror variables have finite range.
- structural: This approach uses the structural abstraction approach [CDJR09] to compute the abstraction. It assumes all the predicates and mirror variables have finite range.
- hybrid: This approach uses the hybrid BDD+SMT abstraction approach \(\left[\mathrm{CCF}^{+} 07, \mathrm{CFG}^{+} 10\right]\) to compute the abstraction. It assumes all the predicates and mirror variables have finite range.
- bdd: This approach uses the BDD existential quantification to compute the abstraction. It assumes all the predicates and mirror variables have finite range. Moreover, it assumes the input model to be finite-state (i.e. it must not contain neither real nor integer variables).

Specifies the refinement strategy to be used within the CEGAR loop to refine the abstraction when the abstract counter-example is spurious. Possible values for this variable are:
- itp: (default) It preforms the refinement analyzing the interpolants [McM03] induced by the spurious counter-example. The counter-example is split in two parts by considering each state of the counterexample. The first part of the formula consists of the prefix of the counter-example from initial state to the considered state (included). The second part is the suffix of the counter-example from the considered state (included) till the last state of the counter-example. The interpolants will be on the variables corresponding to the considered states (i.e. on the language in the intersection among the two formulas).
- uc: It preforms the refinement analyzing the unsatisfiable core induced by the unsatisfiability in the concrete model of the abstract counter-example. This approach has a limitation that considers only the predicates in the extracted unsatisfiability core that do not refer to variables at different states in the counter-example. (The predicate \(@(a, 0)=@(b, 2)\) where variable \(a\) is at state 0 and variable \(b\) at state 2 will be discarded. While \(@(a, 0)=@(b, 0)\) will be considered. Both variable refer to the same state.)
- wp: It preforms the refinement computing weakest preconditions induced by the spurious abstract counter-example.

\section*{msat_dump_format}

Environment Variable
This variable controls the format used by commands like e.g. msat_check_invar_bmc when option \(-d\) is given to the command. The valid values for this variable are:
- mathsat: (default) This format is the language specific of the MATHSAT [CGSS13] SMT solver.
- smtlib: This format is standard format [BST12] adopted in the SMT competition and accepted by (almost) all the SMT solvers at the state-of-the-art.

This is a Boolean variable that specifies the format used while printing counter-examples or while writing infinite precision Real constants. Its default value is 0 , meaning that for instance \(f^{\prime} 2 / 3\) is used to represent the rational number \(2 / 3\). If set to 1 , then the output would be 0.6666666667 .

This is a Boolean variable that specifies whether when interacting with the SMT solver to perform upfront bit-blasting or to pass the words directly to the SMT solver. By default this variable is set to 1 , meaning that the words are passed natively to the SMT solver.

\section*{qe.engine}

Environment Variable
Specifies the high level quantifier elimination engine to be used while computing the abstraction during e.g. the CEGAR loop. Possible values for this variable are:
- msat: This approach uses AllSMT [LNO06] to compute the abstraction. It assumes all the predicates and mirror variables have finite range.
- structural: This approach uses the structural abstraction approach [CDJR09] to compute the abstraction. It assumes all the predicates and mirror variables have finite range.
- hybrid: (default) This approach uses the hybrid BDD+SMT abstraction approach \(\left[\mathrm{CCF}^{+} 07\right.\), \(\mathrm{CFG}^{+} 10\) ] to compute the abstraction. It assumes all the predicates and mirror variables have finite range.

\section*{qe.hybrid.backjumping_enabled}

Environment Variable
This is a Boolean variable that enables the back-jumping optimization within the TCC encoder [CCF \({ }^{+} 07\), \(\mathrm{CFG}^{+} 10\) ] when the qe.engine is set to hybrid. By default this variable is set to 0 , i.e. the optimization is disabled.

This is a Boolean variable that enables the D'Agostino optimization when the qe.engine is set to hybrid. By default this variable is set to 0 , i.e. the optimization is disabled.

This is a Boolean variable that enables the conjunctive partitioning of the formula to abstract when the qe.engine is set to hybrid. By default this variable is set to 0 , i.e. the optimization is disabled.

This is a Boolean variable that enables the use of a threshold for computing the conjunctive partitioning of the formula to abstract when the qe.engine is set to hybrid. By default this variable is set to 0 , i.e. the optimization is disabled.

\section*{qe.hybrid.threshold_value}

\section*{Environment Variable}

This is an positive integer variable that specifies the maximm size in terms of BDD nodes for each conjuct of the formula to abstract when the conjunctive partitioning is enabled and the qe.engine is set to hybrid. By default this variable is set to 300 BDD nodes.

This variable controls the low level engine used when the qe.engine is set to msat or structural. Possible values are:
- allsmt: (default) This value specifies to use AllSMT approach. It assumes that all the infinite domain variables (i.e. real and integer) are quantified out.
- fm: This value specifies to use Fourier-Motzking [Sch98] quantifier elimination technique. It assumes the model contains real variables and no integer.
- 1 w : This value specifies to use Loos-Weispfenning [LW93, Mon08] quantifier elimination technique. It assumes the model contains real variables and no integer.
```

qe.msat.remove_redundant_constraints_enabled
Environment Variable

```

This is a Boolean variable that enables the optimization that pre-process the formula to remove redundant part of the formula within the SMT solver. This option takes effect when the qe.engine is set to msat and the qe.msat.engine is set to 1 w or \(f \mathrm{~m}\). By default this variable is set to 0 , i.e. the optimization is enabled.

\section*{qe.msat.boolean_simplifications_enabled} Environment Variable

This is a Boolean variable that enables the optimization that enables the Boolean simplification of the formula within the SMT solver. This option takes effect when the qe.engine is set to msat and the qe.msat.engine is set to \(l w\) or \(f m\). By default this variable is set to 0 , i.e. the optimization is enabled.

This is a Boolean variable that enables the optimization within the SMT solver to push quantifiers inside the formula. This option takes effect when the qe.engine is set to msat and the qe.msat.engine is set to 1 w or \(f m\). By default this variable is set to 0 , i.e. the optimization is enabled.

\section*{qe.structural.analyze_conjuncts_enabled}

Environment Variable
This is a Boolean variable that enables the analysis of the possible conjuncts of the formula to abstract to see whether pushing of quantifiers could be performed. This option takes effect when the qe.engine is set to structural. By default this variable is set to 0 , i.e. the optimization is disabled.

This is a Boolean variable that enables the assertion in the SMT solver of all the partial results of quantification of the conjuncts of the formula to abstract while performing the quantification. This option takes effect when the qe.engine is set to structural. By default this variable is set to 0 , i.e. the optimization is disabled.

This variable specifies the core engine used to perform quantifier elimination when the qe.engine is set to structural. By default this variable is set to hybrid, i.e. the engine used is the BDD+SMT approach of \(\left[\mathrm{CCF}^{+} 07, \mathrm{CFG}^{+} 10\right]\).

\section*{qe.structural.dagostino_enabled}

Environment Variable
This is a Boolean variable that enables the D'Agostino optimization when the qe.engine is set to structural. By default this variable is set to 0 , i.e. the optimization is disabled.

This is a Boolean variable that enables the conversion of the formula to quantify in Disjunctive Normal Form (DBF) when the qe.engine is set to structural. By default this variable is set to 0 , i.e. the optimization is disabled.

This is a Boolean variable that enables the generation of intermediate BDDs for each leaf (quantifier free formula resulting from the quantifier elimination) element of the formula to quantify when the qe.engine is set to structural. By default this variable is set to 0 , i.e. the optimization is disabled.

\section*{qe.structural.incrementality_enabled}

Environment Variable
This is a Boolean variable that enables the exploitation of the incrementality of the SMT solver while performing the quantification of the formula to abstract when the qe.engine is set to structural. By default this variable is set to 0 , i.e the optimization is disabled.

This is a Boolean variable that enables the inlining of equalities while performing the quantification of the formula to abstract when the qe. engine is set to structural. By default this variable is set to 0 , i.e the optimization is disabled.

This variable affects the behavior of the code that performs the inlining of expressions. In particular it specifies the number of iterations to perform to discover possible equivalences to perform the inlining of conjuncts. Default value is 0 , i.e. perform a fix-point.

This is a Boolean variable that enables the low level quantifier optimizations [CDJR09] while performing the quantification of the formula to abstract when the qe.engine is set to structural. By default this variable is set to 1 , i.e. the optimization is enabled.

This is a Boolean variable that enables the pre-assertion in the SMT solver of all the conjuncts, if any, of the formula to abstract when the qe.engine is set to structural. By default this variable is set to 0 , i.e the optimization is disabled.

This is a Boolean variable that enables the variable sampling optimization [CDJR09] while performing the quantification of the formula to abstract when the qe.engine is set to structural. By default this variable is set to 1 , i.e. the optimization is enabled.

\section*{write xmi max word_width}

Environment Variable
This variable controls the maximum number of bits allowed for the bit vectors when dumping the model in XMI format with the command write_xmi_model. The default value is 6 .

Remark: Large values may lead to huge times in dumping the XMI format. Indeed, for each value of the word an XMI state may be created.

\section*{Chapter 6}

\section*{Running NUXMV batch}
nuXmv so far provides an batch interaction inherited from the original NuSMV. We report here the different command line options provided both by NuSMV and by nuXmv.

When the -int option is not specified, nuXmv runs as a batch program, in the style of SMV, performing (some of) the steps described in previous section in a fixed sequence.
```

system_prompt> nuXmv [command line options] input-file <RET>

```

The program described in input-file is processed, and the corresponding finite state machine is built. Then, if input-file contains formulas to verify, their truth in the specified structure is evaluated. For each formula which is not true a counterexample is printed.
The batch mode can be controlled with the following command line options:
```

nuXmv [-h | -help] [-v vl] [-int]
[[-source script_file | -load script_file]]
[-s] [-old] [-old_div_op] [-smv_old]
[-disable_syntactic_checks]
[-keep_single_value_vars]
[-disable_daggifier] [-dcx] [-cpp] [-pre pps]
[-ofm fm_file] [-obm bm_file] [-lp]
[-n idx] [-is] [-ic] [-ils] [-ips] [-ii] [-ctt]
[[-f] [-r]]|[-df] [-flt] [-AG]
[-coi] [-i iv_file] [-o ov_file]
[-t tv-file] [-reorder] [-dynamic] [-m method]
[-disable_sexp2bdd_caching] [-bdd_soh heuristics]
[[-mono]|[-thresh cp_t]|[-cp cp_t]|[-iwls95 cp_t]]
[-noaffinity] [-iwls95preorder]
[-bmc] [-bmc_length k]
[-sat_solver name] [-sin on|off] [-rin on|off]
[-ojeba algorithm] [ input-file]

```
where the meaning of the options is described below. If input-file is not provided in batch mode, then the model is read from standard input.
\(\underset{\substack{\text { Fovedzione } \\ \text { BRINO KESSER }}}{ }\)
\begin{tabular}{ll}
-help \\
-h & \\
-v verbose-level & \begin{tabular}{l} 
Prints the command line help. \\
Enables printing of additional information on the internal operations of \\
nuXmv. Setting verbose-level to 1 gives the basic information. Using this \\
option makes you feel better, since otherwise the program prints nothing un- \\
til it finishes, and there is no evidence that it is doing anything at all. Setting \\
the verbose-level higher than 1 enables printing of much extra information.
\end{tabular} \\
Enables interactive mode
\end{tabular} of pre-processor names contained within double quotes.
\(\underset{\substack{\text { Fovedzione } \\ \text { BRINO KESSER }}}{ }\)
\begin{tabular}{|c|c|}
\hline -ofm fm-file & Prints flattened model to file fn_file \\
\hline -obm bm_file & Prints boolean model to file bn_file \\
\hline -lp & Lists all properties in SMV model \\
\hline -n idx & Specifies which property of SMV model should be checked \\
\hline -is & Does not check SPEC properties. Sets to " 1 " the ignore_spec environment variable. \\
\hline -ic & Does not check COMPUTE properties. Sets to " 1 " the ignore_compute environment variable. \\
\hline -ils & Does not check LTLSPEC properties. Sets to " 1 " the ignore_ltlspec environment variable. \\
\hline -ips & Does not check PSLSPEC properties. Sets to " 1 " the ignore_pslspec environment variable. \\
\hline -ii & Does not check InVARSPEC properties. Sets to "1" the ignore_invariant environment variable. \\
\hline -ctt & Checks whether the transition relation is total. \\
\hline -f & Computes the set of reachable states before evaluating CTL expressions. Since NuSMV-2.4.0 this option is set by default, and it is provided for backward compatibility only. See also option -df. \\
\hline -r & Prints the number of reachable states before exiting. If the -f option is not used, the set of reachable states is computed. \\
\hline -df & Disable the computation of the set of reachable states. This option is provided since NuSMV-2.4.0 to prevent the computation of reachable states that are otherwise computed by default. \\
\hline -flt & Forces the computation of the set of reachable states for the tableau resulting from BDD-based LTL model checking (command check_ltlspec). If the option -flt is not specified (default), the resulting tableau will inherit the computation of the reachable states from the model, if enabled. If the option \(-f l t\) is specified, the reachable states set will be calculated for the model and for the tableau resulting from LTL model checking. This might improve performances of the command check_ltlspec, but may also lead to a dramatic slowing down. This options has effect only when the calculation of reachable states is enabled (see -f). \\
\hline -AG & Verifies only AG formulas using an ad hoc algorithm (see documentation for the ag_only_search environment variable). \\
\hline -coi & Enables cone of influence reduction. Sets to " 1 " the cone_of_influence environment variable. We remark that, when cone of influence reduction is enabled, a counter-example trace for a property that does not hold may not be a valid counter-example trace for the original model. We refer the reader to the Frequently Asked Questions (FAQ) [FAQ]. \\
\hline
\end{tabular}
\begin{tabular}{|c|c|}
\hline -i iv-file & Reads the variable ordering from file iv file. \\
\hline - o ov_file & Writes the variable ordering to file \(o v_{-}\)file. \\
\hline -t tv file & Reads a variable list from file \(t v_{-}\)file. This list defines the order for clustering the transition relation. This feature has been provided by Wendy Johnston, University of Queensland. The results of Johnston's et al. research have been presented at FM 2006 in Hamilton, Canada. See [WJKWLvdBR06]. \\
\hline -reorder & Enables variable reordering after having checked all the specification if any. \\
\hline -dynamic & Enables dynamic reordering of variables \\
\hline -m method & Uses method when variable ordering is enabled. Possible values for method are those allowed for the reorder_method environment variable (see the NUSMV user manual [CCCJ \(\left.{ }^{+} 10\right]\) ). \\
\hline \multicolumn{2}{|l|}{-disable_sexp2bdd_caching} \\
\hline & Sets the default value of environment variable enable_bdd_cache to 0 , i.e. the evaluation of symbolic expression to ADD and BDD representations are not cached. See command clean_sexp2bdd_cache for reasons of why BDD cache should be disabled sometimes. \\
\hline -bdd_soh heuristics & Sets the default value of environment variable bdd_static_order_heuristics to heuristics, i.e. the option sets up the heuristics to be used to compute BDD ordering statically by analyzing the input model. See the documentation about variable bdd_static_order_heuristics in the NUSMV user manual [CCCJ \({ }^{+}\)10] for more details. \\
\hline -mono & Enables monolithic transition relation \\
\hline -thresh cp_t & conjunctive partitioning with threshold of each partition set to \(c p \_t\) (DEFAULT, with \(c p_{-} t=1000\) ) \\
\hline -cp cp_t & DEPRECATED: use thresh instead. \\
\hline -iwls95 cp_t & Enables Iwls95 conjunctive partitioning and sets the threshold of each partition to \(c p_{-} t\) \\
\hline -noaffinity & Disables affinity clustering \\
\hline -iwls95preoder & Enables Iwls95CP preordering \\
\hline -bmc & Enables BMC instead of BDD model checking (works only for LTL properties and PSL properties that can be translated into LTL) \\
\hline -bmc_length \(k\) & Sets bmc_length variable, used by BMC \\
\hline -sat_solver name & Sets sat_solver variable, used by BMC so select the sat solver to be used. \\
\hline -sin on,off & Enables (on) or disables (off) Sexp inlining, by setting system variable sexp_inlining. Default value is off. \\
\hline -rin on,off & Enables (on) or disables (off) RBC inlining, by setting system variable rbc_inlining. Default value is on. The idea about inlining was taken from [ABE00] by Parosh Aziz Abdulla, Per Bjesse and Niklas Eén. \\
\hline -ojeba algorithm & Sets the algorthim used for BDD-based language emptiness of Büchi fair transition systems by setting system variable oreg_justice_emptiness_bdd_algorithm (default is EL_bwd). The available algorithms are: EL bwd EL fwd \\
\hline
\end{tabular}

\footnotetext{
The available algorithms are: EL_bwd EL_fwd
}

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\section*{Appendix A}

\section*{Typing and Production Rules}

This appendix contains the syntactic production rules for writing a nuXmv program.

\section*{Identifiers}
```

identifier ::
identifier_first_character
| identifier identifier_consecutive_character
identifier_first_character :: one of
A B C D E F G H I J K L M N O P Q R S T U V W X Y Z

```

```

identifier_consecutive_character ::
identifier_first_character
digit
one of \$ \# -
digit :: one of 0}10

```

Note that there are certain reserved keyword which cannot be used as identifiers (see page 7).

\section*{Variable and DEFINE Identifiers}
```

define_identifier :: complex_identifier
variable_identifier :: complex_identifier

```

\section*{Complex Identifiers}
```

complex_identifier ::
identifier
| complex_identifier . identifier
complex_identifier [ simple_expression ]
self

```

\section*{Integer Numbers}
```

integer_number ::
pos_integer_number
| - pos_integer_number
pos_integer_number ::
digit

```
| pos_integer_number digit

\section*{Real Numbers}
```

real_number ::
float_number
| fractional_number
| exponential_number
float_number ::
pos_integer_number . pos_integer_number
fractional_number ::
fraction_prefix ' pos_integer_number / pos_integer_number
fraction_prefix ::
one of f F
exponential_number ::
pos_integer_number exponential_prefix integer_number
| float_number exponential_prefix integer_number
exponential_prefix ::
one of e E

```

\section*{Constants}
```

constant ::
boolean_constant
| integer_constant
| real_constant
symbolic_constant
word_constant
range_constant
boolean_constant :: one of
FALSE TRUE
integer_constant :: integer_number
real_constant :: real_number
symbolic_constant :: identifier
word_constant :: 0 [word_sign_specifier] word_base [word_width] _ word_value
word_sign_specifier :: one of
u s
word_width :: integer_number (>0)
word_base :: b | B | O | O | d | D | h | H
word_value ::
hex_digit
| word_value hex_digit
| word_value -
hex_digit :: one of
O 1 2 3 4 5 6 7 8 9 a b c def A B C D E F

```

FONDAZIONE

Note that there are some additional restrictions on the exact format of word constants (see page 12).
```

range_constant ::
integer_number .. integer_number

```

\section*{Basic Expressions}
```

basic_expr ::
constant -- a constant
| variable_identifier -- a variable identifier
| define_identifier -- a define identifier
| ( basic_expr )
| ! basic_expr -- logical/bitwise NOT
| basic_expr \& basic_expr -- logical/bitwise AND
| basic_expr | basic_expr -- logical/bitwise OR
| basic_expr xor basic_expr -- logical/bitwise exclusive OR
| basic_expr xnor basic_expr -- logical/bitwise NOT xor
| basic_expr -> basic_expr -- logical/bitwise implication
| basic_expr <-> basic_expr -- logical/bitwise equivalence
| basic_expr = basic_expr -- equality
| basic_expr != basic_expr -- inequality
| basic_expr < basic_expr -- less than
| basic_expr > basic_expr -- greater than
| basic_expr <= basic_expr -- less than or equal
| basic_expr >= basic_expr -- greater than or equal
| - basic_expr -- unary minus
| basic_expr + basic_expr -- integer addition
| basic_expr - basic_expr -- integer subtraction
| basic_expr * basic_expr -- integer multiplication
| basic_expr / basic_expr -- integer division
| basic_expr mod basic_expr -- integer remainder
| basic_expr >> basic_expr -- bit shift right
| basic_expr << basic_expr -- bit shift left
| basic_expr [ index ] -- index subscript
| basic_expr [ integer_number : integer_number ]
-- word bits selection
basic_expr :: basic_expr -- word concatenation
word1 ( basic_expr )
-- boolean to word[1] convertion
bool ( basic_expr )
-- word[1] and integer to boolean convertion
toint ( basic_expr )
-- word[N] and boolean to integer convertion
signed ( basic_expr )
-- unsigned to signed word convertion
unsigned ( basic_expr )
-- signed to unsigned word convertion
extend ( basic_expr , basic_expr)
-- word width increase
resize ( basic_expr , basic_expr)
-- word width resizing
| basic_expr union basic_expr
-- union of set expressions
{ set_body_expr } -- set expression
basic_expr in basic_expr -- inclusion expression
basic_expr ? basic_expr : basic_expr
-- if-then-else expression
count ( basic_expr_list )
-- count of TRUE boolean expressions
floor ( basic_expr )
case_expr -- case expression
next ( basic_expr ) -- next expression
basic_expr_list ::
basic_expr
| basic_expr_list , basic_expr
set_body_expr ::

```
\(\underset{\substack{\text { Fonvazione } \\ \text { BRINO KESS } \\ \hline}}{ }\)
```

basic_expr
| set_body_expr , basic_expr

```

\section*{Case Expression and If-Then-Else Expression}
```

case_expr :: case case_body esac
case_body ::
basic_expr : basic_expr ;
| case_body basic_expr : basic_expr ;
basic_expr ? basic_expr : basic_expr

```

\section*{Simple Expression}
```

simple_expr :: basic_expr

```

Note that simple expressions cannot contain next operators.

\section*{Next Expression}
```

next_expr :: basic_expr

```

\section*{Type Specifier}
```

type_specifier ::
simple_type_specifier
| module_type_spicifier
simple_type_specifier ::
boolean
word [ integer_number ]
unsigned word [ integer_number ]
signed word [ integer_number ]
integer
real
{ enumeration_type_body }
integer_number .. integer_number
array integer_number .. integer_number
of simple_type_specifier
enumeration_type_body ::
enumeration_type_value
| enumeration_type_body , enumeration_type_value
enumeration_type_value ::
symbolic_constant
| integer_number

```

\section*{Module Type Specifier}
```

module_type_specifier ::
identifier [ ( [ parameter_list ] ) ]
% | process identifier [ ( [ parameter_list ] ) ]
parameter_list ::
simple_expr
| parameter_list , simple_expr

```

Fonoizione

\section*{State, Input and Frozen Variables}
```

var_declaration :: VAR var_list
ivar_declaration :: IVAR simple_var_list
frozenvar_declaration :: FROZENVAR simple_var_list
var_list :: identifier : type_specifier ;
| var_list identifier : type_specifier ;
simple_var_list :: identifier : simple_type_specifier ;
| simple_var_list identifier : simple_type_specifier ;

```

\section*{DEFINE Declaration}
```

define_declaration :: DEFINE define_body
define_body :: identifier := simple_expr ;
| define_body identifier := simple_expr ;

```

\section*{CONSTANTS Declaration}
```

constants_declaration :: CONSTANTS constants_body ;
constants_body :: identifier
| constants_body , identifier

```

\section*{ASSIGN Declaration}
```

assign_constraint :: ASSIGN assign_list
assign_list :: assign ;
| assign_list assign ;
assign ::
complex_identifier := simple_expr
| init ( complex_identifier ) := simple_expr
| next ( complex_identifier ) := next_expr

```

\section*{TRANS Statement}
```

trans_constraint :: TRANS next_expr [;]

```

\section*{INIT Statement}
```

init_constrain :: INIT simple_expr [;]

```

\section*{INVAR Statement}
```

invar_constraint :: INVAR simple_expr [;]

```

\section*{Module Declarations}
```

module :: MODULE identifier [(module_parameters)] [module_body]
module_parameters ::
identifier
| module_parameters , identifier

```
\(\underset{\substack{\text { Fonvazione } \\ \text { BRINOK KESS ER }}}{ }\)
```

module_body ::
module_element
| module_body module_element
module_element ::
var_declaration
| ivar_declaration
| frozenvar_declaration
| define_declaration
| constants_declaration
| assign_constraint
| trans_constraint
| init_constraint
| invar_constraint
| fairness_constraint
| ctl_specification
| invar_specification
| ltl_specification
| compute_specification
| isa_declaration

```

\section*{ISA Declaration}
```

isa_declaration :: ISA identifier

```

Warning: this is a deprecated feature and will eventually be removed from NUSMV. Use module instances instead.

\section*{CTL Specification}
```

ctl_specification :: SPEC ctl_expr ;
ctl_expr ::
simple_expr -- a simple boolean expression
| ( ctl_expr )
| ! ctl_expr -- logical not
| ctl_expr \& ctl_expr -- logical and
| ctl_expr | ctl_expr -- logical or
| ctl_expr xor ctl_expr -- logical exclusive or
| ctl_expr xnor ctl_expr -- logical NOT exclusive or
| ctl_expr -> ctl_expr -- logical implies
| ctl_expr <-> ctl_expr -- logical equivalence
EG ctl_expr -- exists globally
EX ctl_expr -- exists next state
EF ctl_expr -- exists finally
AG ctl_expr -- forall globally
AX ctl_expr -- forall next state
AF ctl_expr -- forall finally
E [ ctl_expr U ctl_expr ] -- exists until
A [ ctl_expr U ctl_expr ] -- forall until

```

\section*{INVAR Specification}
```

invar_specification :: INVARSPEC simple_expr ;

```

This is equivalent to
```

SPEC AG simple_expr ;

```
but is checked by a specialised algorithm during reachability analysis.


\section*{LTL Specification}
```

ltl_specification :: LTLSPEC ltl_expr [;]
ltl_expr ::
simple_expr -- a simple boolean expression
| ( ltl_expr )
| ! ltl_expr -- logical not
| ltl_expr \& ltl_expr -- logical and
| ltl_expr | ltl_expr -- logical or
| ltl_expr xor ltl_expr -- logical exclusive or
| ltl_expr xnor ltl_expr -- logical NOT exclusive or
| ltl_expr -> ltl_expr -- logical implies
| ltl_expr <-> ltl_expr -- logical equivalence
-- FUTURE
| x ltl_expr -- next state
| G ltl_expr -- globally
| F ltl_expr -- finally
| ltl_expr U ltl_expr -- until
| ltl_expr v ltl_expr -- releases
-- PAST
| Y ltl_expr -- previous state
| Z ltl_expr -- not previous state not
| H ltl_expr -- historically
| O ltl_expr -- once
| ltl_expr S ltl_expr -- since
| ltl_expr T ltl_expr -- triggered

```

\section*{Real Time CTL Specification}
```

rtctl_specification :: SPEC rtctl_expr [;]
rtctl_expr ::
ctl_expr
EBF range rtctl_expr
ABF range rtctl_expr
EBG range rtctl_expr
ABG range rtctl_expr
A [ rtctl_expr BU range rtctl_expr ]
| E [ rtctl_expr BU range rtctl_expr ]
range :: integer_number .. integer_number

```

It is also possible to compute quantative information for the FSM:
```

compute_specification :: COMPUTE compute_expr [;]
compute_expr :: MIN [ rtctl_expr , rtctl_expr ]
MAX [ rtctl_expr , rtctl_expr ]

```

\section*{PSL Specification}
```

pslspec_declaration :: "PSLSPEC " psl_expr ";"
psl_expr ::
psl_primary_expr
| psl_unary_expr
| psl_binary_expr
| psl_conditional_expr
| psl_case_expr
| psl_property

```
\(\underset{\text { FoNDazione }}{\text { BRUNO KESSLER }}\)
```

psl_primary_expr ::
number ;; a numeric constant
boolean ;; a boolean constant
var_id ;; a variable identifier
{ psl_expr , ... , psl_expr }
{ psl_expr "{" psl_expr , ... , "psl_expr" }}
| ( psl_expr )
psl_unary_expr ::
+ psl_primary_expr
| - psl_primary_expr
| ! psl_primary_expr
psl_binary_expr ::
psl_expr + psl_expr
psl_expr union psl_expr
psl_expr in psl_expr
psl_expr - psl_expr
psl_expr * psl_expr
psl_expr / psl_expr
psl_expr % psl_expr
psl_expr == psl_expr
psl_expr != psl_expr
psl_expr < psl_expr
psl_expr <= psl_expr
psl_expr > psl_expr
psl_expr >= psl_expr
psl_expr \& psl_expr
psl_expr | psl_expr
psl_expr xor psl_expr
psl_conditional_expr ::
psl_expr ? psl_expr : psl_expr
psl_case_expr ::
case
psl_expr : psl_expr ;
psl_expr : psl_expr ;
endcase

```

Among the subclasses of psl_expr we depict the class psl_bexpr that will be used in the following to identify purely boolean, i.e. not temporal, expressions.
```

psl_property ::
replicator psl_expr ; ; a replicated property
FL_property abort psl_bexpr
psl_expr <-> psl_expr
psl_expr -> psl_expr
FL_property
| OBE_property
replicator ::
forall var_id [index_range] in value_set :
index_range ::
[ range ]
range ::
low_bound : high_bound
low_bound ::
number
| identifier
high_bound ::
number
identifier
| inf ;; inifite high bound

```

Fondizione
```

value_set ::
{ value_range , ... , value_range }
boolean
value_range ::
psl_expr
| range
FL_property ::
;; PRIMITIVE LTL OPERATORS
X FL_property
X! FL_property
F FL_property
G FL_property
[ FL_property U FL_property ]
[ FL_property W FL_property ]
; SIMPLE TEMPORAL OPERATORS
always FL_property
never FL_property
next FL_property
next! FL_property
eventually! FL_property
FL_property until! FL_property
FL_property until FL_property
FL_property until!_ FL_property
FL_property until_ FL_property
FL_property before! FL_property
FL_property before FL_property
FL_property before!_ FL_property
FL_property before_ FL_property
; EXTENDED NEXT OPERATORS
X [number] ( FL_property )
X! [number] ( FL_property )
next [number] ( FL_property )
next! [number] ( FL_property )
;
next_a [range] ( FL_property )
next_a! [range] ( FL_property )
next_e [range] ( FL_property )
next_e! [range] ( FL_property )
next_event! ( psl_bexpr ) ( FL_property )
next_event ( psl_bexpr ) ( FL_property )
next_event! ( psl_bexpr ) [ number ] ( FL_property )
next_event ( psl_bexpr ) [ number ] ( FL_property )
; ;
next_event_a! ( psl_bexpr ) [psl_expr] ( FL_property )
next_event_a ( psl_bexpr ) [psl_expr] ( FL_property )
next_event_e! ( psl_bexpr ) [psl_expr] ( FL_property )
next_event_e ( psl_bexpr ) [psl_expr] ( FL_property )
OPERATORS ON SERES
sequence ( FL_property )
sequence |-> sequence [!]
sequence |=> sequence [!]
;;
always sequence
G sequence
never sequence
eventually! sequence
;;
within! ( sequence_or_psl_bexpr , psl_bexpr ) sequence

```
```

    within ( sequence_or_psl_bexpr , psl_bexpr ) sequence
    within!_ ( sequence_or_psl__bexpr , psl__bexpr ) sequence
    within_ ( sequence_or_psl_bexpr , psl_bexpr ) sequence
    ;;
whilenot! ( psl_bexpr ) sequence
whilenot ( psl_bexpr ) sequence
whilenot!_ ( psl_bexpr ) sequence
whilenot_ ( psl_bexpr ) sequence
sequence_or_psl_bexpr ::
sequence
| psl__bexpr
sequence ::
{ SERE }
SERE ::
sequence
psl_bexpr
;; COMPOSITION OPERATORS
SERE ; SERE
| SERE : SERE
| SERE \& SERE
| SERE \&\& SERE
| SERE | SERE
;; RegExp QUALIFIERS
SERE [* [count] ]
[* [count] ]
SERE [+]
[+]
;;
psl_bexpr [= count ]
| psl_bexpr [-> count ]
count ::
number
range
OBE_property ::
AX OBE_property
AG OBE_property
AF OBE_property
A [ OBE_property U OBE_property ]
EX OBE_property
EG OBE_property
EF OBE_property
E [ OBE_property U OBE_property ]

```

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[^0]:    ${ }^{1}$ For more information on these operators see the NUSMV user manual [CCCJ ${ }^{+}$10].

[^1]:    ${ }^{2}$ See 2.3.3) for array defines and 2.3.1 for array variables.

[^2]:    ${ }^{3}$ See 2.1.7 for more information about the set types and their counterpart types
    ${ }^{4}$ See 2.1.7 for more information about the set types and their counterpart types

[^3]:    ${ }^{5}$ See 2.1.7 for information on set types and their counterpart types
    ${ }^{6}$ See Section 2.1.8 [Type Order], page 10 for the information on the order of types.

[^4]:    ${ }^{7}$ Similarly to NUSMV, in the current version of NUXMV, compassion constraints are supported only for BDD-based LTL model checking. We plan to add support for compassion constraints also for CTL specifications and in Bounded Model Checking in forthcoming releases of nuXmv.

[^5]:    ${ }^{8}$ This also means that the actual parameters are analyzed in the context of the variable declaration where the module is instantiated, not in the context of the expression where the formal parameter occurs.

[^6]:    ${ }^{9}$ The module main is instantiated with the so called empty identifier which cannot be referenced in a program.

[^7]:    ${ }^{10}$ Note that if the ordering is not provided by a user then NUXMV decides by itself how to order the variables. Two shell variables bdd_static_order_heuristics (see the NUSMV user manual [CCCJ ${ }^{+}$10] for further information) and vars_order_type allow to control the ordering creation.

[^8]:    gen_Itlspec_bmc - Dumps into one or more dimacs files the given LTL specifica-

[^9]:    ${ }^{1}$ To be exact, $M_{1} \subseteq M_{2}$, where $M_{1}$ is the model from which the trace was generated, and $M_{2}$ is the currently loaded, and built, model. Note however, that this may mean that the trace is not valid for the model $M_{2}$.

