

Models of Memristors for SPICE Simulations

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Abstract— Memristors are novel devices which can be used in applications such as memory, logic, analog circuits, and neuromorphic systems. Several memristor technologies have been developed such as ReRAM (Resistive RAM), MRAM (Magnetoresistance RAM), and PCM (Phase Change Memory). To design circuits with memristors, the behavior of the memristor needs to be described by a mathematical model. While the model for memristors should be sufficiently accurate as compared to the behavior of physical devices, the model must also be computationally efficient. Several models for memristors have been proposed – the linear ion drift model, the nonlinear ion drift model, the Simmons tunnel barrier model, and the ThrEshold Adaptive Memristor (TEAM) model. In this paper, the different memristor models are described and a Verilog-A implementation for these models, including the relevant window functions, are presented. These models are suitable for EDA tools such as SPICE.

Index Terms— memristor, memristive systems, SPICE, Verilog-A, TEAM.

I. INTRODUCTION

Memristors are passive two-port elements with variable resistance (also known as a memristance) [1]. Changes in the memristance depend upon the history of the device (e.g., the memristance may depend on the total charge passed through the device, or alternatively, on the integral over time of the applied voltage between the ports of the device). Memristive systems [2] are an extension to memristors, where a current-controlled time-invariant memristive device is represented by

$$\frac{dw}{dt} = f(w, i), \quad (1)$$

$$v(t) = R(w, i) \cdot i(t), \quad (2)$$

where w is an internal state variable, $i(t)$ is the current of the memristive device, $v(t)$ is the voltage across the memristive device, $R(w, i)$ is the memristance, and t is time. The terms memristor and memristive systems are often used interchangeably to describe memristive devices.

Memristors can be used in applications such as memory, logic, analog circuits, and neuromorphic systems. A memristor offers several advantages as compared to standard memory technologies: nonvolatility, good scalability, effectively no leakage current, and compatibility with CMOS technology, both electrically and in terms of manufacturing. Several

memristor technologies have been developed such as ReRAM (Resistive RAM), MRAM (Magnetoresistance RAM), and PCM (Phase Change Memory).

To design circuits with memristors, the behavior of the memristor needs to be described by a mathematical model. While the model for memristors should be sufficiently accurate as compared to the behavior of physical devices, it must also be computationally efficient. It is also desirable for the model to be simple, intuitive, and closed-form, as well as general so that it can be tuned to suit different technologies of memristors. Several memristor models have been proposed: the linear ion drift model, the nonlinear ion drift model, the Simmons tunnel barrier model, and the ThrEshold Adaptive Memristor (TEAM) model. In this paper, the different memristor models are described and a Verilog-A code for these models and the relevant window functions are presented. These models are suitable for EDA tools such as SPICE.

II. MEMRISTOR MODELS

All of the memristor models which have been implemented in the Verilog-A model are described in [3]. In this paper, only a brief description of these models is provided. The basic equations and the main characteristics of the memristor models are listed in Table 1. A user manual to this Verilog-A model is provided in [4].

A. Linear Ion Drift Model

In the linear ion drift model [5], two resistors are connected in series, one resistor represents the high concentration of dopants region (high conductance) and the second resistor represents the oxide region (low conductance). A linear ion drift in a uniform field is also assumed, where the ions have equal average ion mobility μ_v . This model exhibits the definition of the original memristor in [1], but is inaccurate as compared to physical memristive devices.

B. Nonlinear Ion Drift Model

In the nonlinear ion drift model [6], a voltage-controlled memristor exhibiting a nonlinear dependence between the voltage and the internal state derivative is assumed. In this model, the state variable w is a normalized parameter within the interval $[0, 1]$. This model also assumes an asymmetric switching behavior.

C. Simmons Tunnel Barrier Model

The Simmons tunnel barrier model [7] assumes nonlinear and asymmetric switching behavior due to an exponential dependence of the movement of the ionized dopants, namely, changes in the internal state variable. In this model, rather than two resistors in series as in the linear drift model, there is a resistor in series with an electron tunnel barrier. The state variable x is the Simmons tunnel barrier width.

D. Threshold Adaptive Memristor (TEAM) Model

The TEAM model [3] is a general memristor model. In this model, a current threshold and tunable nonlinear (polynomial) dependence between the current and the derivative of the internal state variable are assumed. The current-voltage relationship can be in a linear or an exponential manner. It is possible to fit the TEAM model to the Simmons tunnel barrier model or to any different memristor model and gain a more efficient computational time with sufficient accuracy.

III. WINDOW FUNCTIONS

To maintain the physical bounds of the device and add nonlinear behavior close to these physical bounds, several window functions are implemented in the Verilog-A model. These window functions are: Jogelkar [8], Biolek [9], Prodrumakis [10], and TEAM (named Kvatinsky in the Verilog-A model). The window functions and main properties are listed in Table 2.

IV. VERILOG-A CODE

```

`include "disciplines.vams"
`include "constants.h"

// define meter units for w parameter nature distance
access = Metr;
units = "m";
abstol = 0.01n;
endnature
discipline Distance
potential distance;
enddiscipline

module Memristor(p, n, w_position);
input p; // positive pin
output n; // negative pin
output w_position; // w-width pin
electrical p, n, gnd;
Distance w_position;
ground gnd;
parameter real model = 0; // define the model 0 - Linear
// Ion Drift ; 1 - Simmons Tunnel Barrier; 2 - Team model;
// 3- Nonlinear Ion Drift model
parameter real window_type=0; // define the window type :
// 0 - No window; 1 - Jogelkar window ; 2 - Biolek window ;
// 3 - Prodrumakis window ; 4- Kvatinsky window (Team
// model only)
parameter real dt=0; // user must specify dt same as max step

```

```

// size in transient analysis & must be at least 3 orders
// smaller than T period of the source
parameter real init_state=0.5; // the initial state condition
// [0:1]

```

```

/////////////////////////////////Linear Ion Drift model/////////////////////////////////
//parameters definitions and default values for linear model
parameter real Roff = 200000;
parameter real Ron = 100;
parameter real D = 3n;
parameter real uv = 1e-15;
parameter real w_multiplied = 1e8; // transformation factor
// for w/X width in meter units
parameter real p_coeff = 2; // Windowing function
// coefficient
parameter real J = 1; // for prodromakis Window function
parameter real p_window_noise=1e-18; // provoke the w
// width not to get stuck at 0 or D with p window
parameter real treshhold_voltage=0;

```

```

// local variables
real w;
real dwdt;
real w_last;
real R;
real sign_multiply;
real stp_multiply;
real first_iteration;

```

```

///////////////////////////////// Simmons Tunnel Barrier model/////////////////////////////////
//parameters definitions and default values
parameter real f_off = 3.5e-6;
parameter real f_on = 40e-6;
parameter real i_off = 115e-6;
parameter real i_on = 8.9e-6;
parameter real x_c = 107e-12;
parameter real b = 500e-6;
parameter real a_on = 2e-9;
parameter real a_off = 1.2e-9;

```

```

// local variables
real x;
real dxdt;
real x_last;

```

```

/////////////////////////////////TEAM model/////////////////////////////////
parameter real K_on=-8e-13;
parameter real K_off=8e-13;
parameter real Alpha_on=3;
parameter real Alpha_off=3;
parameter real IV_relation=0; // IV_relation=0 means linear
// V=IR. IV_relation=1 means nonlinear V=I*exp{..}
parameter real x_on=0;
parameter real x_off=3e-09; // equals D
// local variables
real lambda;

```

```

////////////////////////////////Nonlinear Ion Drift model////////////////////////////////
parameter real alpha = 2;
parameter real beta = 9;
parameter real c = 0.01;
parameter real g = 4;
parameter real N = 14;
parameter real q = 13;
parameter real a = 4;

analog function integer sign; //Sign function for Constant
// edge cases
real arg; input arg;
sign = (arg >= 0 ? 1 : -1);
endfunction
analog function integer stp; //Stp function
real arg; input arg;
stp = (arg >= 0 ? 1 : 0);
endfunction

////////////////////////////////MAIN //////////////////////////////////
analog begin
if(first_iteration==0) begin
w_last=init_state*D; // if this is the first iteration,
// start with w_init
x_last=init_state*D; // if this is the first
// iteration, start with x_init
end

////////////////////////////////Linear Ion Drift model //////////////////////////////////
if (model==0) begin // Linear Ion Drift model
dwdt =(uv*Ron/D)*I(p,n);
//change the w width only if the threshold_voltage permits!
if(abs(I(p,n))<treshhold_voltage/R) begin
w=w_last;
dwdt=0;
end
if ((window_type==0)|| (window_type==4)) begin // No
// window
w=dwdt*dt+w_last;
end // No window
if (window_type==1) begin // Jogelkar window
if (sign(I(p,n))==1) begin
sign_multiply=0;
if(w==0) begin
sign_multiply=1;
end
end
if (sign(I(p,n))==-1) begin
sign_multiply=0;
if(w==D) begin
sign_multiply=-1;
end
end
end
w=dwdt*dt*(1-pow(2*w/D-
1,2*p_coeff))+w_last+sign_multiply*p_window_noise;
end // Jogelkar window

if (window_type==2) begin // Biolek window
if (stp(-I(p,n))==1) begin
stp_multiply=1;
end
if (stp(-I(p,n))==0) begin
stp_multiply=0;
end
w=dwdt*dt*(1-pow(w/D-
stp_multiply,2*p_coeff))+w_last;
end // Biolek window
if (window_type==3) begin // Prodromakis window
if (sign(I(p,n))==1) begin
sign_multiply=0;
if(w==0) begin
sign_multiply=1;
end
end
if (sign(I(p,n))==-1) begin
sign_multiply=0;
if(w==D) begin
sign_multiply=-1;
end
end
w=dwdt*dt*J*(1-pow(pow(w/D-
0.5,2)+0.75,p_coeff))+ w_last + sign_multiply *
p_window_noise;
end // Prodromakis window
if (w>=D) begin
w=D;
dwdt=0;
end
if (w<=0) begin
w=0;
dwdt=0;
end

//update the output ports(pins)
R=Ron*w/D+Roff*(1-w/D);
w_last=w;
Metr(w_position) <+ w*w_multiplied;
V(p,n) <+ (Ron*w/D+Roff*(1-w/D))*I(p,n);
first_iteration=1;
end // end Linear Ion Drift model

////////////////////////////////Simmons Tunnel Barrier model////////////////////////////////
if (model==1) begin // Simmons Tunnel Barrier model
if (sign(I(p,n))==1) begin
dxdt =f_off*sinh(I(p,n)/i_off)*exp(-exp((x_last-
a_off)/x_c-abs(I(p,n)/b))-x_last/x_c);
end
if (sign(I(p,n))==-1) begin
dxdt = f_on*sinh(I(p,n)/i_on)*exp(-exp((a_on-
x_last)/x_c-abs(I(p,n)/b))-x_last/x_c);
end
x=x_last+dt*dxdt;
if (x>=D) begin
x=D;

```

```

        dxdt=0;
    end
    if (x<=0) begin
        x=0;
        dxdt=0;
    end
    //update the output ports(pins)
    R=Ron*(1-x/D)+Roff*x/D;
    x_last=x;
    Metr(w_position) <+ x/D;
    V(p,n) <+ (Ron*(1-x/D)+Roff*x/D)*I(p,n);
    first_iteration=1;
end // end Simmons Tunnel Barrier model

////////////////////////////////TEAM model////////////////////////////////
if (model==2) begin // Team model
    if (I(p,n) >= i_off) begin
        dxdt =K_off*pow((I(p,n)/i_off-1),Alpha_off);
    end
    if (I(p,n) <= i_on) begin
        dxdt =K_on*pow((I(p,n)/i_on-1),Alpha_on);
    end
    if ((i_on<I(p,n)) && (I(p,n)<i_off)) begin
        dxdt=0;
    end
    if (window_type==0) begin // No window
        x=x_last+dt*dxdt;
    end // No window
    if (window_type==1) begin // Jogelkar window
        x=x_last+dt*dxdt*(1-pow((2*x_last/D-1),(2*p_coeff)));
    end // Jogelkar window
    if (window_type==2) begin // Biolek window
        if (stp(-I(p,n))==1) begin
            stp_multiply=1;
        end
        if (stp(-I(p,n))==0) begin
            stp_multiply=0;
        end
        x=x_last+dt*dxdt*(1-pow((x_last/D-stp_multiply),(2*p_coeff)));
    end // Biolek window
    if (window_type==3) begin // Prodromakis window
        x=x_last+dt*dxdt*J*(1-pow((pow((x_last/D-0.5),2)+0.75),p_coeff));
    end // Prodromakis window
    if (window_type==4) begin //Kvatinsky window
        if (I(p,n) >= 0) begin
            x=x_last+dt*dxdt*exp(-exp((x_last-a_off)/x_c));
        end
        if (I(p,n) < 0) begin
            x = x_last+dt*dxdt*exp(-exp((a_on-x_last)/x_c));
        end
    end // Kvatinsky window
    if (x>=D) begin

```

```

        dxdt=0;
        x=D;
    end
    if (x<=0) begin
        dxdt=0;
        x=0;
    end
    lambda = ln(Roff/Ron);
    //update the output ports(pins)
    x_last=x;
    Metr(w_position) <+ x/D;
    if (IV_relation==1) begin
        V(p,n) <+ Ron*I(p,n)*exp(lambda*(x-x_on)/(x_off-x_on));
    end
    else if (IV_relation==0) begin
        V(p,n) <+ (Roff*x/D+Ron*(1-x/D))*I(p,n);
    end
    first_iteration=1;
end // end TEAM model

////////////////////////////////Nonlinear Ion Drift model////////////////////////////////
if (model==3) begin // Nonlinear Ion Drift model
    if (first_iteration==0) begin
        w_last=init_state;
    end
    dwdt = a*pow(V(p,n),q);
    if ((window_type==0) || (window_type==4)) begin
        // No window
        w=w_last+dt*dwdt;
    end // No window
    if (window_type==1) begin // Jogelkar window
        w=w_last+dt*dwdt*(1-pow((2*w_last-1), (2 * p_coeff)));
    end // Jogelkar window
    if (window_type==2) begin // Biolek window
        if (stp(-V(p,n))==1) begin
            stp_multiply=1;
        end
        if (stp(-V(p,n))==0) begin
            stp_multiply=0;
        end
        w=w_last+dt*dwdt*(1-pow((w_last-stp_multiply),(2*p_coeff)));
    end // Biolek window
    if (window_type==3) begin // Prodromakis window
        w=w_last+dt*dwdt*J*(1-pow((pow((w_last-0.5),2)+0.75),p_coeff));
    end // Prodromakis window
    if (w>=1) begin
        w=1;
        dwdt=0;
    end
    if (w<=0) begin
        w=0;
        dwdt=0;
    end
end

```

```

//change the w width only if the threshold_voltage permits!
if(abs(V(p,n))<treshhold_voltage) begin
w=w_last;
end

//update the output ports(pins)
w_last=w;
Metr(w_position) <+ w;
I(p,n) <+ pow(w,N) *beta *sinh(alpha*V(p,n)) +c*
(exp(g*V(p,n))-1);
first_iteration=1;
end // end Nonlinear Ion Drift model
end // end analog
endmodule

```

V. CONCLUSIONS

A Verilog-A code that contains several models, useful for design in memristor-based circuits, is presented in this paper, as well as relevant window functions. This Verilog-A model can be used by circuit designers, since it is easy to use, contains several mathematical models, the parameters of already existing models can be easily changed, as well as additional mathematical models can be added.

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TABLE 1. THE DIFFERENT MEMRISTOR MODELS (FURTHER DESCRIPTION IN [2])

Model	Linear ion drift [5]	Nonlinear ion drift [6]	Simmons tunneling barrier [7]	TEAM [3]
State variable	$0 \leq w \leq D$ Doped region physical width	$0 \leq w \leq 1$ Doped region normalized width	$a_{off} \leq x \leq a_{on}$ Undoped region width	$x_{on} \leq x \leq x_{off}$ Undoped region width
Control mechanism	Current controlled	Voltage controlled	Current controlled	Current controlled
Threshold	None	None	None	i_{on}, i_{off}

TABLE 2. COMPARISON OF DIFFERENT WINDOW FUNCTIONS

Function	Jogelkar [8]	Biolek [9]	Prodromakis [10]	TEAM [3]
$f(x)/f(w)$	$f(w) = 1 - (2w/D - 1)^{2p}$	$f(w) = 1 - (w/D - stp(-i))^{2p}$	$f(w) = j(1 - [(w - 0.5)^2 + 0.75]^p)$	$f_{on,off} = \exp[-\exp(x - x_{on,off} /w_c)]$
Fits memristor model	Linear/nonlinear ion drift/TEAM	Linear/nonlinear ion drift/TEAM	Linear/nonlinear ion drift/TEAM	TEAM for Simmons tunneling barrier fitting