# TECHNICAL RESEARCH REPORT

A Comparative Study of Reactor Designs for the Production of Graded Films with Applications to Combinatorial CVD

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A Comparative Study of Reactor Designs for the Production of Graded Filmswith Applications to Combinatorial CVD

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#### Abstract:

Segmented CVD reactor designs enabling spatial control of across-wafer gas phase composition were evaluated for depositing graded films suitable for combinatorial studies. Specifically two reactor designs were constructed and evaluated with experiments and response surface model (RSM) based analysis to quantify the reactor performance in terms of film thickness uniformity, sensitivity to adjustable reactor operating conditions, range of thickness over which uniformity could be achieved and each reactor's ability to control the thickness gradient across the wafer surface. Design features distinguishing the two reactor systems and their influence on gradient control versus deposition rate performance are summarized. RS models relating wafer state properties to process recipes are shown to be effective tools to quantify, qualify and compare different reactor designs.

#### I. Introduction

The sem iconductor industry constantly innovates and improves process and tool designs in an effort to keep up with Moore's Law. Chemical vapor deposition (CVD) tools are prevalent in every sem iconductor fabrication facility as an efficient method for depositing nonvolatile solid films with good film conformality. However, conventional CVD systems are designed for a narrow range of operating conditions and do not offer much

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flexibility for in proving process recipes and optim izing process development cycles for new materials. Also, most designs do not allow for controlling precursor concentration gradients over a wafer surface during a deposition run allowing for combinatorial capabilities.

There are relatively few examples of chemical vapor deposition reactor systems designed com binatorial capabilities. Those that do exist, how ever, demonstrate the capability to produce films with graded properties over a portion of the substrate surface. For example, the CVD reactor design of Gladfelter [1-2] features three feed tubes in a triangular arrangement across the substrate; a different single-source precursor is fed through each tube, generating compositional spreads of three metal dioxides over the substrate. In Wang [3-5], thickness graded films of hydrogenated silicon were deposited in a hot-wire CVD system featuring a mask and motorized shutter; control of the shutter spead was used to create strips of graded films over the substrate. Finally, in Taylor and Semancik [6], microhotplate devices were used to control the temperature in an array of micro-scale substrate samples; it was found that tem perature quadients in the m icrohotplate supports resulted m icrostructurally graded film on the support legs.

Earlier work [7-9] by the authors of this study describes the prelim inary construction and testing of a spatially program mable chemical vapor deposition (SP-CVD) system that was developed at the University of Maryland. The original SP-CVD reactor design henceforth denoted as design A), construction, operation and preliminary evaluation experiments are described in the cited references. Figure 1 depicts a schematic diagram of design A comprising the individually controllable segmented showerhead with segments S1, S2 and S3 arranged over the wafer surface. For this and the previous studies, we consider blanket tungsten by  $H_2$  reduction of W  $F_6$  as the model deposition system; the overall deposition reaction is:

 $W F_6 (q) + 3H_2 (q) \rightarrow W (s) + 6HF (q)$ .

The results from the earlier work cited demonstrated for the first time the SP-CVD system's ability to be reprogrammed, effectively reconfiguring the reactor solely in software between deposition runs to intentionally induce spatially non-uniform thickness deposition patterns on a single wafer. In [1], a relatively simple linearmodel was used to

relate average film thickness under each of the segments to the feed gas recipe of each segment. Because this model did not account for segment to segment interactions, a more accurate modeling approach is developed in this paper to enable modeling of those interactions. The purpose of this paper is to demonstrate the use of Response Surface Models (RS models) to predict film thickness response over the entire wafer to adjustable process parameters enabling control to a specified thickness spatial function, such as a linear thickness gradient across a patch of wafer surface. This model is used to quantify the reactor's performance and examine the relative merits of different reactor designs. This approach is applied to evaluate two reactor designs: the "original" SP-CVD reactor (design A) and a modification (design B) motivated by an attempt to reduce the chamber volume.

#### III.M odeling for design A

Key to this study is the development of an accurate model of the fullwafer response to adjustable process operating conditions; the model is necessary to compute process recipes that optimize a wafer profile objective function. The model, while physically motivated, will be identified from a set of experiments.

### A. The response surface approach:

The response surface modeling approach comprises of the following three steps [10]:

- 1) System attic experim ents: This step entails setting up a series of experim ents that generate a range of reliable measurements of the desired output or response variable. The input variables/predictor variables are varied systematically to generate the range of measurements of the response variable by running experiments on the process tool. For the reactor designs discussed in this paper, we selected a subset of experiments based on our intuitive understanding of segment to segment interactions based on the results from preliminary experiments in [1], followed by a statistical analysis of the estimated parameters.
- 2) Identify a mathematical model relating the response variables (wafer thickness profile) to the input variables. The model form (linear vs. quadratic for example) is based on our physical and intuitive understanding of the process. The model is

- tested for accuracy and validated. The derivation of the RS  $\mathfrak m$  odel is discussed in detail the next section.
- 3) The RS model is used to optim ize the settings of the input variables to m inim ize the value of an objective function, based on our film gradient control criterion, solving a constrained non-linear optim ization problem. This optim ization is discussed in section IV, topic C in this paper.

In this paper, the input variables of the reactor system are defined by the recipe of the SP-CVD tool. This recipe comprises the flow rate of  $H_2$ , the flow rate of  $W_6$  ( $H_2:W_6$  flow ratio is fixed at 4:1), and the show exhead-w afergap size. The desired response variable is the film thickness of deposited tungsten defined at a specific spatial resolution over the w afer surface.

#### B.Derivation of the model form:

Under isotherm alprocessing conditions, the overall reaction rate can be expressed as the following surface reaction expression [11]:

$$R_{kin} = k_{o} \left[ P_{WF_{6}} \right]^{0} \left[ P_{H_{2}} \right]^{1/2}$$
 (1)

where,

 $R_{kin}$  is the rate of deposition of tungsten

 $[\![P_{_{W\ F_{_{6}}}}]\!]$  is the partial pressure of W  $\,F_{6}$ 

 $[P_{H_0}]$  is the partial pressure of H  $_2$ 

A coording to this reaction kinetics model, the reaction rate does not depend on W  $F_6$  partial pressure when sufficient W  $F_6$  is present. However, the reaction rate is assumed to be proportional to the square root of the hydrogen precursor concentration  $X_{H_2}$ , so as a first order approximation we have:

$$X_{H_2} = \frac{P_{H_2}}{\text{Total pressure P}} \approx \frac{\text{flow of H}_2 \text{ (sccm.)}}{\text{Total flow of precursor (sccm.)}}$$

Furtherm one, in our experiments we should expect a linear relationship between the deposition rate of the W film and  $\sqrt{X_{\rm H_2}}$  when precursor conversion rates are low, and so the square root of H<sub>2</sub> flow to each segment is finally used as input to our model.

The SP-CVD reactor has a show exhead with three segments which interact with one another by the following two gas transportmechanisms:

- (1) Intersegn ent gap diffusion: In this mechanism, process gases diffuse from one segment to the other segments though the gap between the wafer surface and the bottom of the segments owing to the concentration gradients between the segments when different recipes are used in neighboring segments.
- (2) Intersegnment back diffusion: In this mechanism, process gases diffuse from the common exhaust volume (CEV) back into the segments owing to gas composition differences between the CEV and individual segments; these differences are attributable to different precursor recipes in the different segments or depletion at high deposition rates.

The showerhead-wafer gap is a process parameter that controls segment to segment interaction in the gap region and is included in the RS model. We derive a model which will predict the entire wafer film thickness profile W $^{pred}$  (r, $\theta$ ) (in nm, for a fixed deposition time) based on the following input variables:

$$x_i : \sqrt{H_2 \text{ flow (seem)}}$$
 to segmenti

g:wafer-showerheadgap,mm

Based on the deposition rate expression, and the mechanism of segment to segment interaction through inter-segment gap diffusion and back diffusion from CEV, we intuitively define the properties of the model to satisfy the following requirements:

- 1) The model should be such that it predicts the local thickness under segment i to be proportional to  $x_i$  and, to a lesser extent  $x_j$  for  $i \neq j$  because of back diffusion.
- 2) Segment i film thickness dependency on  $x_j$  is modulated by g for transport to that region by intersegment gap diffusion.
- 3) No deposition should take place when all  $x_{i=0}$  and the deposition rate should not change with g alone.

Under these assumptions, we arrive at the RS model form to be:

$$W_{\text{pred}} (\mathbf{r}, \theta) = b_{1} (\mathbf{r}, \theta) x_{1} + b_{2} (\mathbf{r}, \theta) x_{2} + b_{3} (\mathbf{r}, \theta) x_{3}$$

$$+ b_{14} (\mathbf{r}, \theta) x_{1} g + b_{24} (\mathbf{r}, \theta) x_{2} g + b_{34} (\mathbf{r}, \theta) x_{3} g$$
(2)

To understand this model, consider a spatial point  $(r',\theta')$  under segment 1; the terms in the expression  $b_1(r,\theta)x_1 + b_2(r,\theta)x_2 + b_3(r,\theta)x_3$  are 'designed' to satisfy requirement 1, i.e.,  $W_{pred}(r',\theta')$  would be primarily dependent on  $x_1$  and the coefficient  $b_1$  quantifies

this dependency. The terms  $b_2(\mathbf{r}',\theta')\mathbf{x}_2$  and  $b_3(\mathbf{r}',\theta')\mathbf{x}_3$  account for the contribution of back diffusion to the point thickness W  $_{\mathrm{pred}}(\mathbf{r}',\theta')$ .

The terms in the expression  $b_{1,A}(r,\theta)x_1g+b_{2,A}(r,\theta)x_2g+b_{3,A}(r,\theta)x_3g$  are designed to satisfy requirement 2, i.e. if the spatial point  $(r',\theta')$  is under segment 1,  $W_{pred}(r',\theta')$  also will depend on the intersegment gap diffusion which is captured by the terms  $b_{1,A}(r',\theta')x_1g,b_{2,A}(r',\theta')x_2g$ , and  $b_{3,A}(r',\theta')x_3g$ . Finally, the absence of a constant term in equation (2) satisfies requirement 3, i.e. when  $x_i=0$  and  $y_i=0$ ,  $y_i=0$ .

The six spatially varying coefficients  $b_i(r,\theta)$  and  $b_{i,j}(r,\theta)$  are computed from the solution of the least squares procedure using the N experimentally determined thickness maps and corresponding process recipes, where N>6. The unique computational approach necessary to compute the spatially varying coefficients will be discussed in a separate publication.

# C.Data set to build RSM for design A:

25 w afers were processed for creating the data set from which we derived the RS model. Each wafer was dipped into 10% HF solution to remove native silicon-oxide film and impurities that block the nucleation of tungsten crystals; after cleaning, the wafers were immediately loaded onto the substrate heater in the reaction chamber. For all experiments described in this article, the heater temperature is set at 400C giving an approximate wafer temperature of 380C. Deposition time was 900 seconds for all wafers. All experiments were carried out at a reactor pressure of 1 torr maintained by a downstream throttle valve.

Table 1 sum marizes a set of experiments which were carried out to generate films of varying thicknesses under different segments by varying the flow rates of the precursor gases and the showerhead-wafer gap sizes. A fter each deposition process, film thickness was measured using a 4 point probe (4PP) ex-situ metrology station. The 4-point probe measurements result in a rectangular grid of measurements over the wafer surfaces with an approximate spatial resolution of 3.45 mm generating 900 measurement points. Numerical analysis of these wafer maps begins by interpolating the thickness data to a numerical quadrature grid defined on a computational domain that has the same physical dimensions as the wafer (see e.g. [12] for the underlying numerical methods, and [13] for

another CVD application). This quadrature grid also is used for num erical interpolation of film thickness in each segment to give a finer (higher resolution) representation of film thickness under each segment.

# D.RSm odelidentification and validation for design A

The six spatially varying coefficients  $b_i(r,\theta)$  and  $b_{i,j}(r,\theta)$  are computed from the solution of the least squares procedure using the 25 experimentally determined thickness maps and corresponding process recipes.

Figure 2 illustrates the comparison between the model's prediction and true measurement for wafers No. 6, 8 and 23 (Table 1). These wafers were processed with the reactor operating in the non-uniformity mode. The RS model was used to predict the segment averaged values which show a good agreement with the true segment averaged values as shown by the bar charts. 10 wafers were processed with the same recipe, operating the reactor in the uniformity mode. This recipe (40 sccm of  $\rm H_2$  in S1 and S3 and 20 sccm of  $\rm H_2$  in S2) was calculated to be the recipe required to produce a thickness of 660 nm in each segment using a linear model. See Ref. 1 for details regarding this linear model. The average thickness of these 10 wafers are calculated and illustrated as a wafermap in Fig. 2 and compared with the RS model's prediction for the same. The RS model predicts the uniformity in agreement with the measured values to an accuracy of 8% with a standard deviation of 5%. Thus RS models can be effectively used to predict the thickness maps accurately produced by the reactor when operated in both the non-uniformity mode and the uniformity mode.

#### IV . Perform ance analysis for design A:

The validated RS model was used to evaluate the original reactor design using the following three criteria:

A . Sensitivity of film thickness profile to gas flow rate and gap: D ifferentiating equation (2) with respect to  $x_1$  we obtain:

$$\frac{\partial W_{\text{pred}}(\mathbf{r},\boldsymbol{\theta})}{\partial \mathbf{x}_{1}} = \mathbf{b}_{1}(\mathbf{r},\boldsymbol{\theta}) + \mathbf{b}_{1,4}(\mathbf{r},\boldsymbol{\theta})\mathbf{g}$$
(3)

Sim ilarly we obtain:

$$\frac{\partial W_{\text{pred}}(\mathbf{r}, \boldsymbol{\theta})}{\partial \mathbf{x}_{2}} = \mathbf{b}_{2}(\mathbf{r}, \boldsymbol{\theta}) + \mathbf{b}_{2,4}(\mathbf{r}, \boldsymbol{\theta}) \mathbf{g} \text{ and}$$
(4)

$$\frac{\partial W_{\text{pred}}(\mathbf{r},\boldsymbol{\theta})}{\partial \mathbf{x}_{3}} = \mathbf{b}_{3}(\mathbf{r},\boldsymbol{\theta}) + \mathbf{b}_{3,4}(\mathbf{r},\boldsymbol{\theta})\mathbf{g}$$
(5)

These sensitivity maps with respect to the reactant concentration of gas fed to each segment are calculated for all  $(r,\theta)$  over the patch of the wafer surface under each segment. Color plots of these patches are found in Fig. 3 showing the sensitivity of W  $_{pred}$   $(r,\theta)$  in each segment to each  $x_i$  with increasing gap size. Two effects are observed here:

- 1) The thickness of film W  $_{pred}$   $(r,\theta)$  in each segment patch i is most sensitive to the corresponding  $x_i$ . This observation is physically intuitive.
- 2) The sensitivity decreases with increasing gap size because as gap size increases, precursor gases 'escape' into the external chamber volume causing reactant depletion over the wafer surface.

Because of the second of these two observations, design A performs poorly when gap sizes are greater than 3 mm. For large gap sizes, the conversion rates are reduced and program m ability of the reactor cannot be exploited for desired uniform ity/non-uniform ity profiles.

# B.R ange of segm ent to segm ent uniform ity

W ith the potential lim itations in operating perform ance for large gap sizes in m ind, the RS m odel is used to predict the range of uniform film thicknesses that can be produced across all segments (i.e., same thicknesses in all three segments) given the fixed range of flow rates allowed by the mass flow controllers (MFCs) for varying gap size. The MFCs for WF<sub>6</sub> have a range from 0 sccm to 12 sccm limiting the  $\rm H_2$  flow range from 0 sccm to 48 sccm. The gap size ranges from 0 to 5 mm.

# Wedefine:

 $W_{i}^{s}$  to be the average thickness for segment i

 $\mathbf{x}_{i}^{s}$  to be the recipe for segment i to achieve W  $_{i}^{s}$ 

 $S_i$  to be the segmentarea, i=1,2,3

We then write out the expression for  $W_i^s$  using (2) and the above defined terms in compactmatrix form as:

$$\begin{bmatrix} W_{1}^{s} \\ W_{2}^{s} \\ W_{3}^{s} \end{bmatrix} = B \begin{bmatrix} x_{1}^{s} \\ x_{2}^{s} \\ x_{3}^{s} \end{bmatrix} + DG \begin{bmatrix} x_{1}^{s} \\ x_{2}^{s} \\ x_{3}^{s} \end{bmatrix}$$

$$(6)$$

where the segm ent-averaged RS model coefficients are

$$B_{i,j} = \int_{S_i} b_j (r_i \theta) ds / \int_{S_i} ds_j; i, j = 1,2,3,$$
(7)

$$D_{i,j} = \int_{S_i} b_{j,4} (r,\theta) ds / \int_{S_i} ds_i; i, j=1,2,3$$
(8)

and the influence of gap size g is included as the m atrix

$$G = \begin{bmatrix} g & 0 & 0 \\ 0 & g & 0 \\ 0 & 0 & g \end{bmatrix}$$
 (9)

To calculate the range of uniform ity that can be achieved using the reactor, we use the following steps:

- 1) Setg=0 m m , W  $^{\rm s} \rightarrow$  0 (desired uniform thickness)
- 2) Set  $W_1^s = W_2^s = W_3^s = W_3^s$
- 3) Solve equation (6) for unknowns  $x_1^{\rm s}$  ,  $x_2^{\rm s}$  and  $x_3^{\rm s}$  .
- 4) If  $x_i^s$  are valid (positive and below the MFC upper limit), then set  $W^s = W^s + \varepsilon_w$  (increment thickness) and return to step 2 i.e., if the unknowns  $x_1^s$ ,  $x_2^s$  and  $x_3^s$  are within the acceptable flow range, we return to step 2, increase the value of the desired uniform thickness and recalculate the unknowns in step 3.
- 5) If  $x_1^s$  are not valid, then  $setg=g+\varepsilon_g$  (increment gap), W  $^s\to 0$  i.e., if the unknowns do not lie between 0 and  $\sqrt{48}$  sccm, we conclude that the desired uniform thickness cannot be achieved with the current gap size and mass flow constraints and return to step 1 and increment the gap size provided it is less than 5 mm.
- 6) If  $g < 5 \, \text{mm}$ , return to step 2.

We thus calculate the maximum value of W  $^{\rm s}$  for a given gap size given the mass flow constraints of the reactor. Figure 4 contains a plot of max W  $^{\rm s}$  vs. gap size. The plot indicates that this reactor can be used to produce uniform films ranging from 0 to 800 nm across all three segments, given the above mentioned range of flow rates. As gap size exceeds  $\sim 3.4$ mm, max W  $^{\rm s}$  reduces to zero. Thus design A is limited to depositing uniform films of thickness < 800 nm and cannot produce segment to segment uniform films for g>3.4 mm. We hypothesize that the curve does not gradually tail off as g grows, but ends abruptly because of segment region asymmetries, such as non-uniform CEV concentration or heater hot spots.

## C. Gradient control perform ance

The program mable reactor can be used to produce wafers with a deliberate thickness gradient across segment regions. To demonstrate this ability, we define a set point film gradient over a subset of the wafer by defining a line of length  $s_n$  on the wafer surface starting at point S1 and ending on S3 (Fig. 5a). The segment wall separating S1 and S3 bisects this line. We define the desired thickness gradient along this line  $W_{\rm set}$  (s) by the following equation:

$$W_{\text{set}}(s) = \frac{2s - s_{\text{m}}}{s_{\text{m}}} W_{\text{m}} \delta_{\text{m}} + W_{\text{avg}}$$
(10)

w here

 $0 \le s \le s_m$ ,  $W_{avg}$  is the mean thickness along the gradient defined by the user in nm, and  $W_m$  is the difference between the values of the thickness at the two extremities of the gradient and the centre point, defined by the user in nm.

 $\delta_{\rm m}$  is a tuning parameter that varies from -1 to 1. This parameter is used so that equation (10) represents all possible linear thickness gradients from S1 to S3 over the length s. Consider, for example,

$$W_{\text{set}}(s) = -\left(\frac{2W_{\text{m}}}{s_{\text{m}}}\right)s + W_{\text{avg}} + W_{\text{m}}) \text{ when } \delta_{\text{m}} = -1 \text{ (m axim um negative gradient)}$$
 (11)

$$W_{set}(s) = W_{avg}$$
 when  $\delta_m = 0$  (flat profile) (12)

$$W_{\text{set}}(s) = \left(\frac{2W_{\text{m}}}{s_{\text{m}}}\right)s + W_{\text{avg}} - W_{\text{m}}) \text{ when } \delta_{\text{m}} = +1 \text{ (m axim um positive gradient)}$$
 (13)

Equation (11) is the equation of a line with negative slope implying that the desired film thickness  $W_{\rm set}$  (s) decreases from S1 to S3 in a linear fashion. Equation (12) is the equation of a horizontal line, implying that the desired film thickness remains constant at  $W_{\rm avg}$  nm from S1 to S3. Finally, equation (13) is an equation of a line with a positive slope, implying that the desired film thickness increases from S1 to S3 in a linear fashion. All other values of  $\delta_{\rm m}$  between -1 and 1 represent the remaining gradients between

$$-(\frac{2W_m}{s_m})$$
 and  $+(\frac{2W_m}{s_m})$  respectively.

In each of the above cases, we set the thickness gradient over any desired target circular patch on the wafer surface, as shown in Fig. 5. The gradient along one axis of the patch is defined by (10), while along the orthogonal axis the gradient is set to zero resulting in a flat tilted circular set point patch,  $W_{\rm set}(r,\theta)$ . Our objective is to calculate a recipe that when inserted into the RS model gives  $W_{\rm pred}(r,\theta)$  that matches  $W_{\rm set}(r,\theta)$  over the entire target patch as accurately as possible. This can be stated as the following optimization problem:

$$\min_{g,x} \left\| W_{pxed}(r,\theta) - W_{set}(r,\theta) \right\|$$
subject to  $0 \le x_i \le 7 \text{ (secon )}^{05} \quad i = 1,2,3$ 

$$0 \le q \le 5mm$$

We define the objective function by numerically computing the norm of the difference between the computed wafer profile in the patch region and the set point; we use the built-in optimization routine 'fmincon' in MATLAB to solve the constrained optimization problem (14). Figures 5b, 5c and 5d illustrate the W  $_{\rm set}$  (s) and W  $_{\rm pred}$  (s) as thickness maps over the patch area when  $\delta_{\rm m}$  equals -1, 0 and 1 respectively. The parameter O  $_{\rm v}$  is the value of the expression  $\|W\|_{\rm pred}$  (s)-W  $_{\rm set}$  (s) at the end of the optimization.

Figure 6a illustrates the gradient across the circular patch extending from S1 to S3 obtained for different values of  $\delta_{\rm m}$ . Plots of the  ${\rm x_i}$ , g, and O  $_{\rm v}$  computed as solutions to (14) as a function of  $\delta_{\rm m}$  are shown in plots 6b and 6c respectively. When  $\delta_{\rm m}$  = -1 our set point corresponds to a film profile that is thickest under S1 and thinnest under S3 on the

defined circular patch. Intuitively, we would predict that  $x_1$  should be higher than  $x_3$ . The optim ization routine computes a recipe which confirms our intuition. Figure 6b illustrates this recipe with  $x_1\sim 5$  (sccm) $^{0.5}$ , with  $x_2\sim 0$  (sccm) $^{0.5}$  and  $x_3\sim 0$  (sccm) $^{0.5}$ . Intuitively, to maintain a steep thickness gradient we would expect to use a small gap size. The optim ization routine arrives at gap size  $\sim 0$  mm (Fig. 6b) to achieve this desired gradient. When  $\delta_m=0$ , our set point corresponds to a film profile that is flat from S1 to S3. Intuitively, we would predict that  $x_1\sim x_3$ . Figure 6b illustrates this recipe with  $x_1$ ,  $x_2$  and  $x_3\sim 2$  (sccm) $^{0.5}$ . However the gap size is 0 mm. Intuitively we would expect a larger gap size for flat profiles, but because design A yields very poor conversion rates with large gap sizes, the RS model used in the optim ization routine 'recommends' a small gap size even for flat profiles.

When  $\delta_m$  = 1, the target film profile is thickest under S3 and thinnest under S1 on the defined circular patch. Intuitively, we would predict that  $x_3$  should be higher than  $x_1$  using a smaller gap size. The optimization routine computes this recipe to be  $x_1 \sim 0.5$  (scam) $^{0.5}$ ,  $x_2 \sim 5$  (scam) $^{0.5}$ , and  $x_3 \sim 4$  (scam) $^{0.5}$  (Fig. 6b) with a gap size of  $\sim 0$  mm.

Thus the RS model was effectively be used to identify recipes to achieve desired thickness gradients on wafer. O  $_{\rm v}$  as low as ~1nm when  $\delta_{\rm m}$  equaled 0, with a mean of 16 nm over all  $\delta_{\rm m}$  was achieved. Roughly speaking, this is approximately 5% error in achieving our setpoint profile, implying a very good gradient control.

# V. The minireactor (Design B)

The analysis using the RS model fordesign A revealed the following draw backs:

- 1) Gap sizes > 3 mm cannot be used because precursor gases escape into the larger cham ber volume resulting in poor conversion rates. Smaller precursor flow rates would further lower the conversion rate. The use of design A for combinatorial A tomic Layer Deposition (ALD), a future research direction, requires minute quantities of precursors pulsed into the reactor. Most of these pulses would escape into the larger chamber volume resulting in a very poor growth rate. This drawback calls for a smaller chamber volume.
- 2) The maximum thickness that can be uniformly deposited in all three segments is ~800 nm. A smaller chamber volume would improve conversion rates and thicker uniform films can be deposited.

Owing to the above drawbacks of design A, it was decided to design, construct and im plem ent a sm aller cham ber (called a m ini cham ber in Figure 7) volum e. In this article, we refer to the SP-CVD reactor with the minicham ber as design B. The minicham ber was constructed from alum inum and comprised of two parts. 1) The main minicham ber and 2) the lid with appropriately shaped holes through which the segments pass. Figure 7 illustrates a schematic of the front view of the design B. The mini chamber seats around the heater and the wafer. The lid rests on small screws drilled horizontally into the segm entwalls 120 degrees apart. The clearance between the outer segm entwalls and the inner walls of the mini chamber is 0.38 mm. After a wafer is transferred to the wafer lifter by a wafer holder from the load lock chamber and lowered onto the heater, the segm ents are low ered. The lid of the m inicham ber then rests on the upper lip of the m ini cham ber while the segments continue to be lowered closer to the wafer. With the lid resting on the minichamber, the segment-wafer gap can be varied from a minimum of 0 mm to a maximum of 10 mm. The minicham bertogether with the lidenclosed the wafer in a cylindrical volume of diameter ~ 106 mm and a height of ~ 105 mm which is considerably smaller than the chamber volume in design A.

# VI.M odeling for design B

#### A.Data set to build RSM for design A:

28 w afers w ere processed to create the data set from which we derived the RS model. Table 2 sum m arizes this data set. Pre-process cleaning, process tem perature, pressure, and post-process m etrology and num erical interpolation techniques remained the same as they were for design A.

# B.RS model identification and validation for design B

The six spatially varying coefficients  $b_i(r,\theta)$  are computed in the same manner as they were computed for design A using the 28 experimentally determined thickness maps and corresponding process recipes.

Figure 8 illustrates the comparison between the model's prediction and true measurement forwafers No2, 11, 13 and 22 (Table 2). These wafers were processed with the reactor operating in the non-uniform mode. The RS model predicts the uniformity in agreement with the measured values to an accuracy of 14% with a standard deviation of 8%. Compared to the design A, the films deposited by design B are 3-4 times thicker. Design

B confines more precursor gases over the wafer surface and improves reactant conversion by as much as 400%. Howeverm odel accuracy appears to be lower than design B.

## VII. Perform ance analysis for design B:

The validated RS model was used to evaluate design B for the same three performance criteria used to evaluate the reactor design A.

#### A . Sensitivity to gap size

The RS m odel captures the sensitivity of the m ini reactor to  $x_i$  and gap through the color plots in Fig.9. The inferences from the plots are:

- 1) As with design A, the thickness of film W  $_{\rm pred}$  (r, $\theta$ ) in each segment is most sensitive to the corresponding x calculated for that segment.
- 2) This sensitivity does not decrease significantly with increasing gap size because the mini chamber in design B confines the gases preventing their escape to the main chamberas in design A. The slight decrease in sensitivity is attributed to the intersegment diffusion that is facilitated by increasing gap size.

# B.Range of segment to segment uniformity

The range of uniform ity that can be achieved using design B was calculated using the sam e procedure used for design A. Figure 10 shows a plot of max W  $^{\rm s}$  vs. gap size for both design A and the design B. The plot indicates that the design B can be used to produce uniform films across all three segments ranging from 0 to 1800 nm, given the earlier mentioned range of flow rates. Design B can thus be used to produce uniform films at a rate 2 to 3 times that of design A.

#### C.G radient control perform ance

We defined and solved the gradient optim ization problem for design B using the same approach used for design A. Figures 11b, 11c and 11d illustrate the W $_{\rm set}$ (s) and W $_{\rm pred}$ (s) when  $\delta_{\rm m}$  equals -1, 0 and 1 respectively while Figure 12 illustrates optim ized profiles and plots between  $\delta_{\rm m}$  vs. x $_{\rm m}$ ,  $\delta_{\rm m}$  vs. gap and  $\delta_{\rm m}$  vs. 0 $_{\rm v}$ .

When  $\delta_{\rm m}=$  -1 the optim ization routine computes a recipe (Fig. 12b) with  $x_1\sim2.5$  (scam) $^{0.5}$ ,  $x_2\sim0$  (scam) $^{0.5}$  and  $x_1\sim0$  (scam) $^{0.5}$ . The optim ization routine recommends a gap size  $\sim0$  mm to achieve this desired gradient.

When  $\delta_{\rm m}=0$ , the optimization routine computes a recipe (Fig. 12b) with  $x_1\sim0.5$  (scam) $^{0.5}$ , with  $x_2\sim1.5$  (scam) $^{0.5}$  and  $x_3\sim0$  (scam) $^{0.5}$  with a gap size  $\sim5$  mm. This confirms our intuition that we would expect a larger gap size for flat profiles.

When  $\delta_m = 1$ , the optim ization routine computes this recipe to be,  $x_1 \sim 0$  (scam) $^{0.5}$ ,  $x_2 \sim 0$  (scam) $^{0.5}$  and  $x_3 \sim 5$  (scam) $^{0.5}$  (Fig. 12b) with a gap size of  $\sim 1$  mm. Comparing Fig.6b with Fig.12b, we see that design B requires  $\sim 50$ % smaller flow rates of precursor gases (defined by  $x_i$ ) than design A for the same thickness gradient.

## VIII.Concluding Remarks

The RS m odeling approach was used successfully to compare the processing capabilities of two CVD reactor designs and to assess their ability to produce controlled graded films over a sub-section of the wafer surface. The following table compares the merits/demerits of the two designs.

Serial	C riteria of com parison	D esign A	Design B
No			
1	RSM odel fidelity reported as		
	$rac{ extsf{W}_{ extsf{predicted}} -  extsf{W}_{ extsf{m} ext{easured}}}{ extsf{W}_{ extsf{m} ext{easured}}}$	8±5%	14±8%
2	Program m ability (Uniform ity/Non	Good	Good
	uniform ity control)		
3	Sensitivity to gap and flow rate	Sensitivity	Exhibits good
		declines rapidly	sensitivity with
		with increasing	increasing g
		g	
4	Range of segment to segment uniformity for	0-800nm with	0-1800nm with
	the given flow constraints	gaps	gaps
		0 <g<3mm< td=""><td>0<g<2.5m m<="" td=""></g<2.5m></td></g<3mm<>	0 <g<2.5m m<="" td=""></g<2.5m>
5	Gradient control	Good	Good

We conclude that design A could be effectively used to deposit uniform and non-uniform films at low gap sizes accurately and with good repeatability. Design B could be used to

deposit thicker uniform and non-uniform films. The gap size could be effectively used as a knob to control intersequent diffusion in the case of design B.

ALD films from binary and ternary systems control film composition by adjusting the pulsing and purging frequencies of the individual precursors. Film compositions can be varied from one wafer to the next using this approach. However, deliberate composition gradient control within a single wafer deposition run has not been demonstrated for ALD. We are currently studying reactor designs for combinatorial ALD that enable gas composition gradient control over the wafer surface to deposit varying compositions over a single wafer. Design B, because of its smaller volume and higher conversion rates could prove useful for this purpose.

## A cknow ledgem ents

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#### Figure Captions

Figure 1: Schem atic of the SP CVD reactor assembly: design A

Figure 2: True wafer maps (data) of wafers No.6, No.8 and No.23 (Table 1), and averaged profiles of 10 wafers processed with the same recipe, obtained from 4 point probe measurements and numerically interpolated in MATLAB are shown in the top row. The interpolated data are compared to the maps predicted by the RS model shown in the second row. The third row compares the average thickness for each segment through bar charts. The recipe is written in the form at:

$$\sqrt{\text{H}_2 \text{ flow (seconn) in S1}}$$
,  $\sqrt{\text{H}_2 \text{ flow (seconn) in S2}}$ ,  $\sqrt{\text{H}_2 \text{ flow (seconn) in S3}}$  gap

Figure 3: Sensitivity of the reactor to recipe and gap size as predicted by the RS model for design A. The greater the redness of the plotw ithin a segment, the more sensitive that segment is to  $H_2$  flow in that segment. As gap size increases, sensitivity to  $H_2$  flow decreases because with increasing gap, the precursor gases 'escape' into the external volume of the chamber.

Figure 4: The range of uniform ity control possible for the design A as predicted by the RS model. This plot conveys that this reactor design could be used to deposit uniform films ranging from 0 to 800 nm using gap sizes ranging from 0 to 3 mm, with WF<sub>6</sub> flow rates ranging from 0 to 12 sccm in each segment (limited by the MFCs) and H<sub>2</sub> flow rates in each segment ranging from 0 to 48 sccm (to maintain the stoichiometric ratio of 1.4 WF<sub>6</sub>H<sub>2</sub>). Arflow in each segment is 60-(H<sub>2</sub> flow + WF<sub>6</sub>flow) sccm.

Figure 5:G radient control for design A for three cases of  $\delta_m$  values (-1,0 and 1). The value of the m in im ized objective function O  $_{\rm v}$  at the end of the optim ization routine is shown below the plots in each case. W  $_{\rm avg}$ , W  $_{\rm m}$  and O  $_{\rm v}$  in nm .

Figure 6: Evaluation of gradient control across segments 1 and 3 as a function of  $\delta_{\rm m}$  using the RS m odel for design A .

Figure 7: Schematic front view of the SP-CVD reactor with the minichamber (design B). When the segments are lowered, the lid is stopped by the wall of the minichamber while the segments continue to be lowered to the desired segment-wafergap. This design renders a chamber with a reduced volume and overcomes draw backs of design A.

Figure 8: True wafer maps (data) of wafers No2, No11, No13 and No22 (Table 2), obtained from 4 point probe measurements and numerically interpolated in MATLAB are

shown in the top row . They are compared with the predicted maps by the RS model for the same shown in the second row. The third row compares the average thickness for each segment through barcharts. The recipe is written in the form at:

$$\left[\sqrt{\text{H}_{2} \text{ flow (secan) in S1}}, \sqrt{\text{H}_{2} \text{ flow (secan) in S2}}, \sqrt{\text{H}_{2} \text{ flow (secan) in S3}}\right]$$

Figure 9: Sensitivity of the reactor to recipe and gap size as predicted by the RS model for design B. The greater the redness of the plot of the plot within a segment the more sensitive that segment is to  $H_2$  flow in that segment. As gap size increases, sensitivity to  $H_2$  flow does not decrease as in design A because the minimizer the gases escaping into the larger chamber volume.

Figure 10: The range of uniform ity control possible for the design A vs. design B as predicted by the RS model. This plot indicates that design B could be used to deposit uniform films ranging from 0 to 1800 nm using gap sizes ranging from 0 to 25 mm, with W  $F_6$  flow rates ranging from 0 to 12 sccm in each segment (limited by the MFCs) and  $H_2$  flow rates in each segment ranging from 0 to 48 sccm (to maintain the stoichiometric ratio of 1.4 W  $F_6$   $H_2$ ). A rflow in each segment is 60-( $H_2$  flow + W  $F_6$  flow) sccm.

Figure 11:G radient control for design B for three cases of  $\delta_m$  values (-1,0 and 1). The value of the m inim ized objective function  $O_v$  at the end of the optim ization routine is shown below the plots in each case. W  $_{\rm avg}$ , W  $_{\rm m}$  and  $O_v$  in nm .

Figure 12: Evaluation of gradient control across segments 1 and 3 as a function of  $\delta_{\rm m}$  using the RS m odel for design B .

# Table Captions

Table1: W afters 1 to 25 were used to deposit films from the above recipes (varying flow rates and show exhead-w after gaps sizes) for generating the data to obtain the RS model for reactor design A. W  $F_6$   $H_2$  flow ratio in each segment is 1.4. A r flow in each segment is 60-( $H_2$  flow + W  $F_6$  flow) scom

Table 2:W afers 1 to 28 were used to deposit films from the above recipes (varying flow rates and show exhead-w afer gaps sizes) for generating the data to obtain the RS model for deign B.W  $F_6$  H $_2$  flow ratio in each segment is 1.4.Ar flow in each segment is 60-(H $_2$  flow + W  $F_6$  flow) sccm

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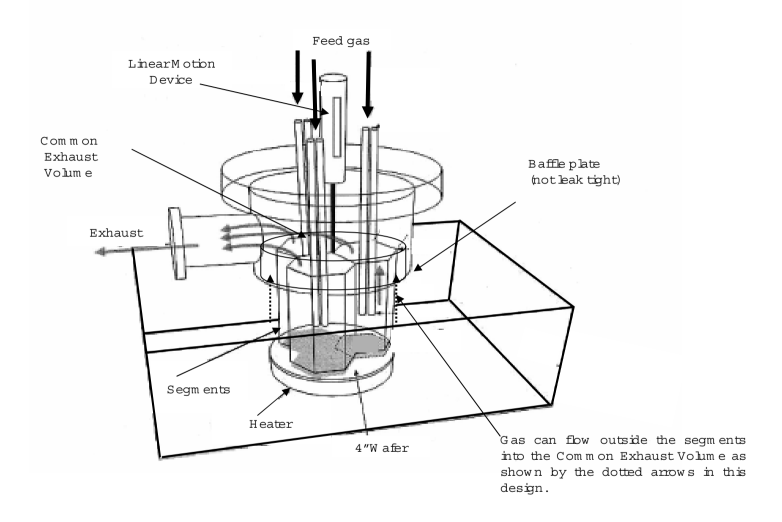


Figure 1

Design A data set

W afernum ber	H2 flow S1(sccm)	H2 flow S2(sccm)	H2 flow S3(sccm)	Gap (mm)
1	24	36	48	1
2	48	24	36	1
3	24	36	48	1
4	48	24	36	1
5	36	48	24	1
6	24	36	48	1
7	48	24	36	1
8	36	48	24	1
9	48	24	36	1
10	40	20	40	1
11	40	20	40	1
12	40	20	40	1
13	40	20	40	1
14	40	20	40	1
15	40	20	40	1
16	40	20	40	1
17	40	20	40	1
18	40	20	40	1
19	40	20	40	1
20	24	36	48	3
21	48	24	36	3
22	36	48	24	3
23	24	36	48	3
24	48	24	36	3
25	36	48	24	3

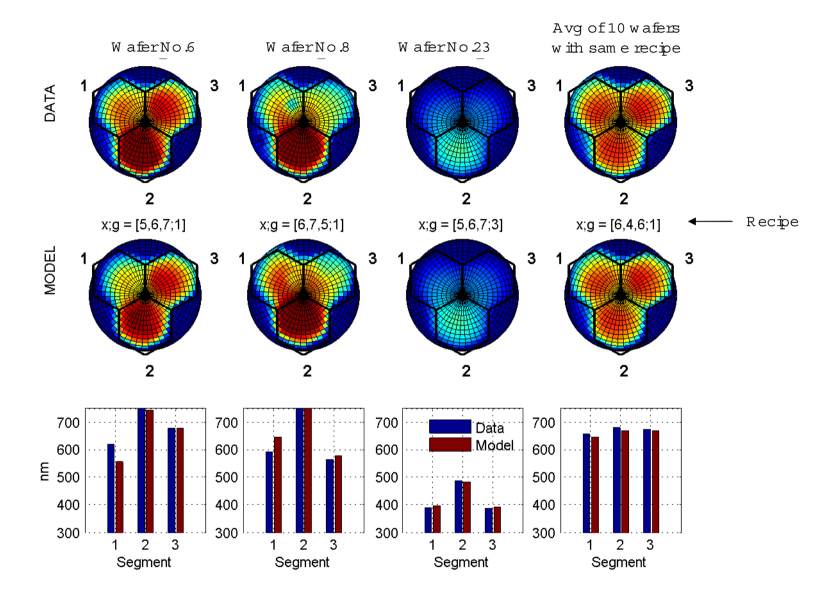
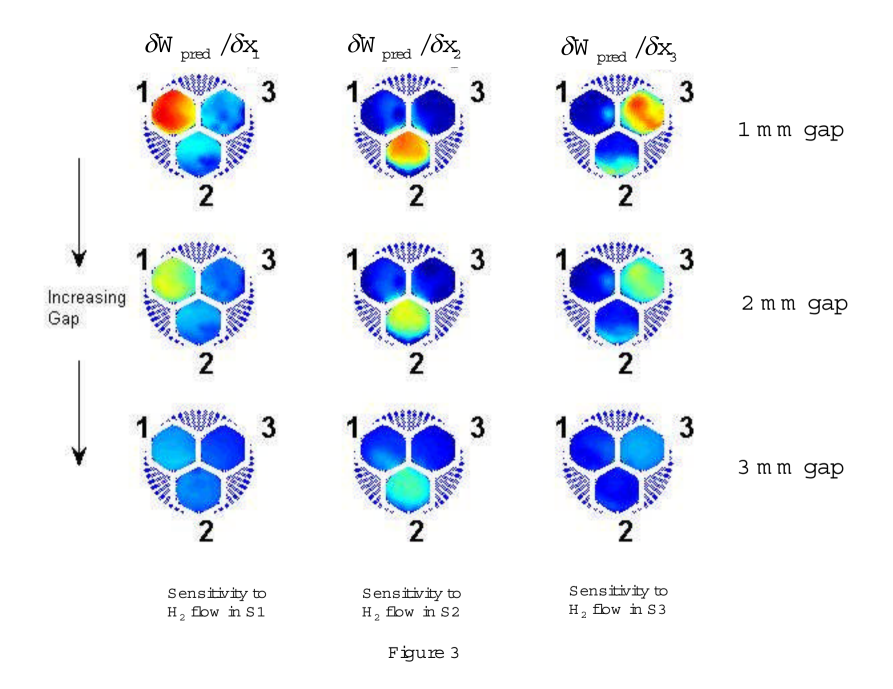


Figure 2



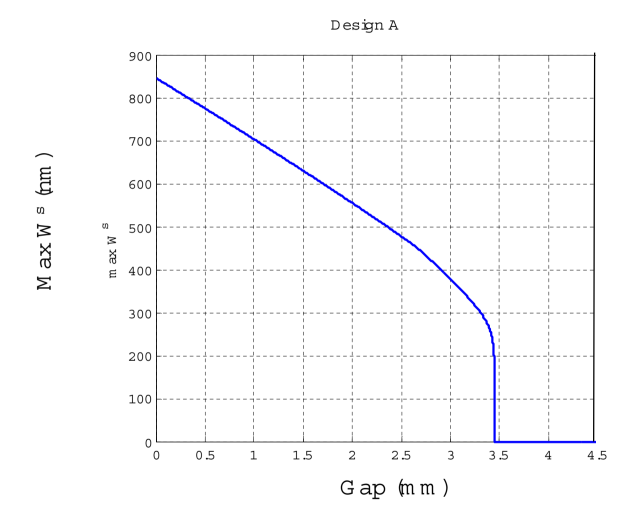


Figure 4

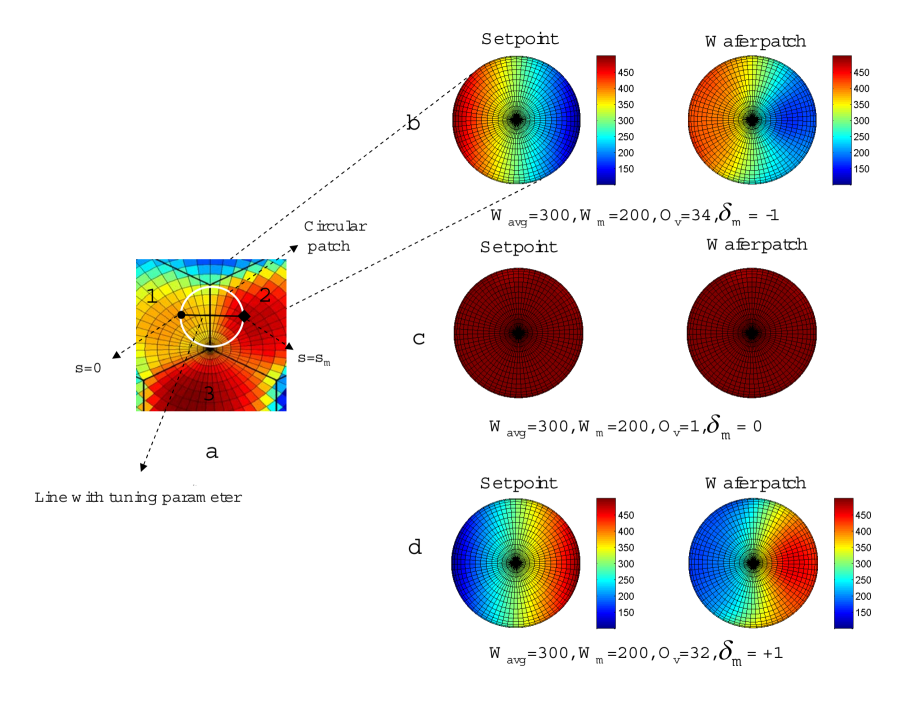


Figure 5

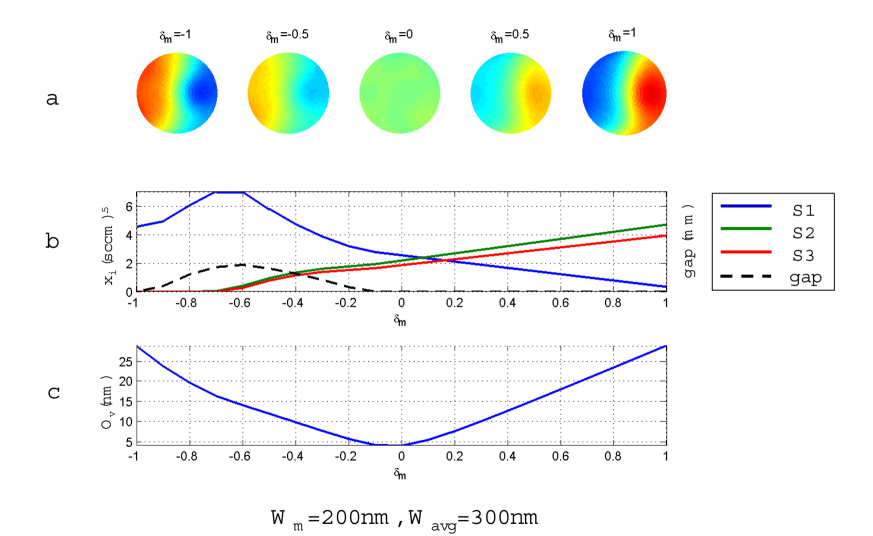


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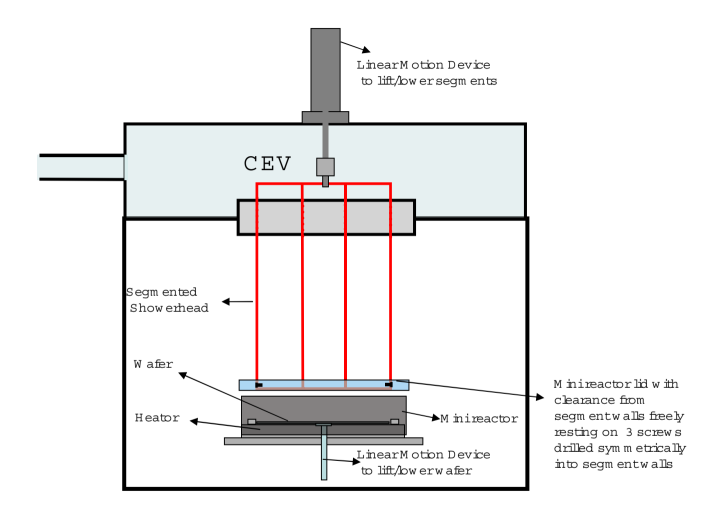


Figure 7

Design B data set

W afernum ber	H2 flow S1 (sccm)	H2 flow S2 (sccm)	H2 flow S3(sccm)	Gap (mm)
1	16	32	48	3
2	48	16	32	3
3	32	32	32	3
4	32	0	0	3
5	0	32	0	3
6	0	0	32	3
7	0	0	32	1
8	0	32	0	1
9	32	0	0	1
10	48	16	32	1
11	32	48	16	1
12	16	32	48	1
13	16	32	48	5
14	32	48	16	5
15	32	32	32	5
16	32	0	0	5
17	0	32	0	5
18	0	0	32	5
19	0	0	32	3
20	0	32	0	3
21	32	0	0	3
22	32	32	32	3
23	48	16	32	3
24	32	48	16	3
25	16	32	48	3
26	0	0	32	2
27	0	32	0	2
28	32	0	0	2

Table 2

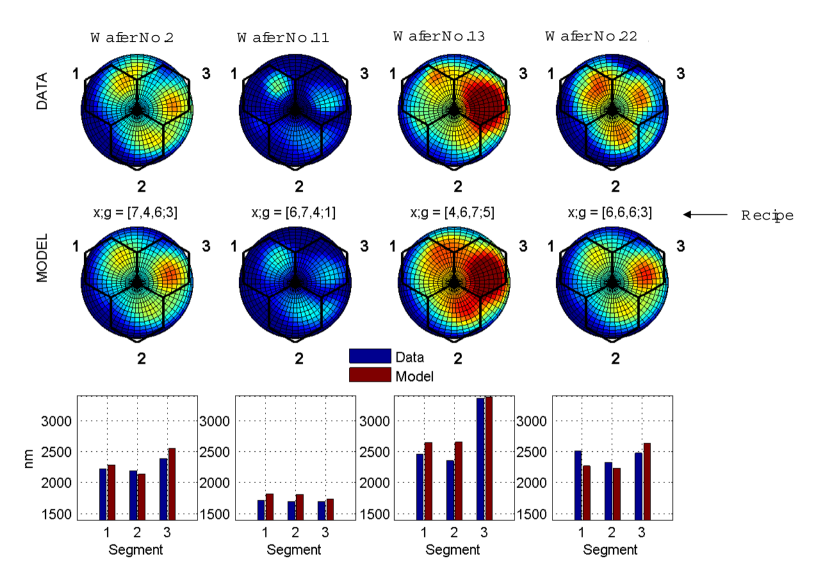


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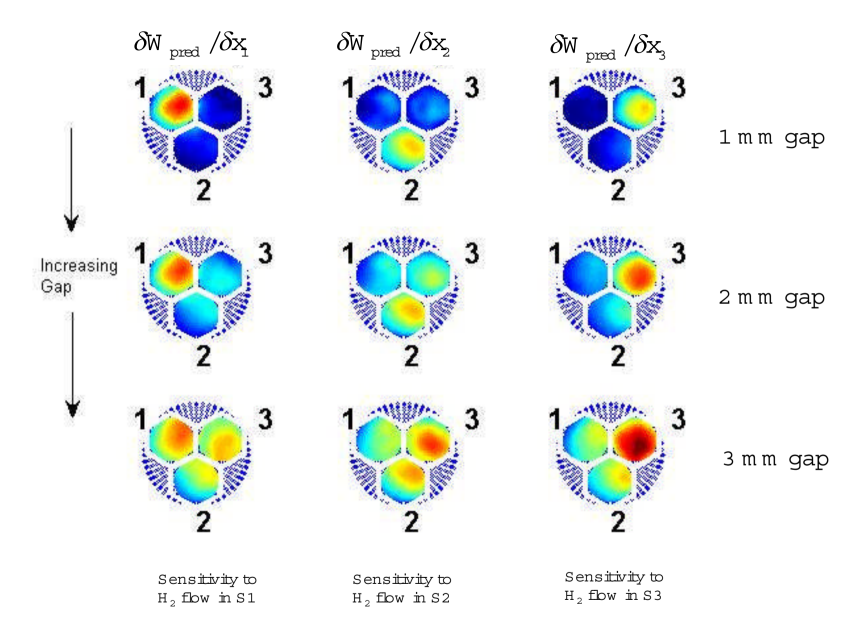


Figure 9

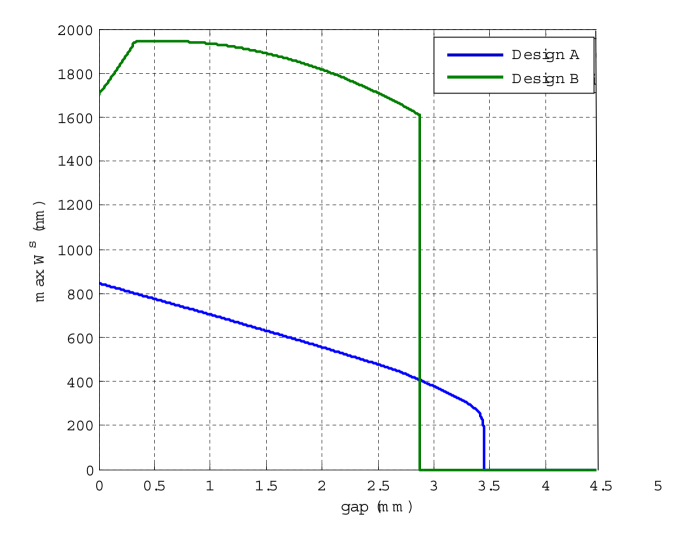


Figure 10

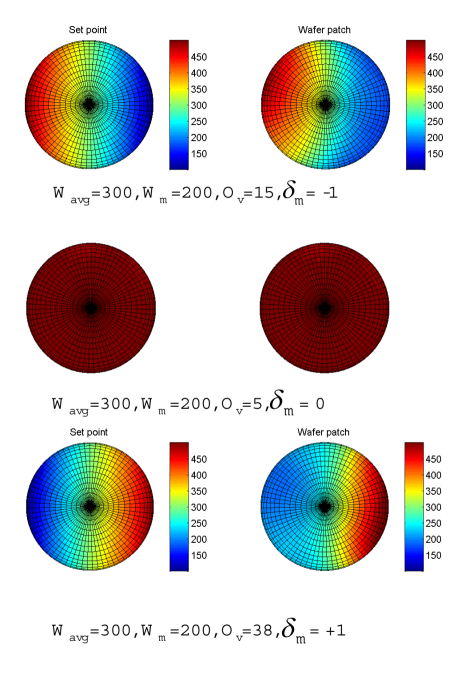


Figure 11

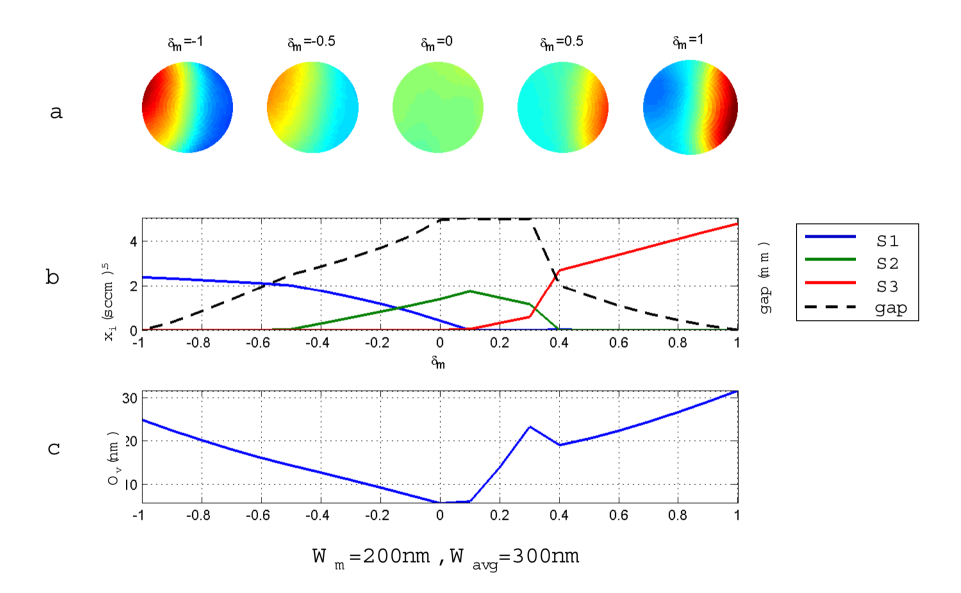


Figure 12