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To cite this version:
This is an author’s version published in: http://oatao.univ-toulouse.fr/20301

Official URL: https://doi.org/10.1016/j.mee.2013.07.022

To cite this version:

Testa, Fabrice and Coetsier, Clémence and Carretier, Emilie and Ennahali, Mohamed and Laborie, Bernard and Moulin, Philippe Recycling a slurry for reuse in chemical mechanical planarization of tungsten wafer: Effect of chemical adjustments and comparison between static and dynamic experiments. (2014) Microelectronic Engineering, 113, 114-122. ISSN 0167-9317

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Recycling a slurry for reuse in chemical mechanical planarization of tungsten wafer: Effect of chemical adjustments and comparison between static and dynamic experiments

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\textbf{A R T I C L E I N F O}

\textbf{Keywords:}
- Tungsten film
- Chemical mechanical polishing
- Slurry
- Chemical adjustments
- Design of experiments

\textbf{A B S T R A C T}

Recycling abrasive slurry that has been used in chemical mechanical polishing (CMP) is one of the options for reducing the cost of manufacturing microchip processors. We use ultrafiltration which is a method of choice to recycle silica (SiO\textsubscript{2})-based slurry. Taking into account that the chemical composition of abrasive slurry plays an important role in tungsten CMP (W-CMP), chemical adjustments have to be made so that the concentrated after used slurry can be reused. In this study, we investigate the effects of chemical additives (iron catalyst, oxalic acid as complexing agent and surfactants as stabilizers) in slurry that has been retreated by ultrafiltration. Experiments are conducted both under static and dynamic conditions and results are compared to better understand the effect of chemical adjustments on the main performances of W-CMP. An optimal chemical adjustment is proposed through a design of experiments evaluation to obtain a concentrated after used and chemically adjust slurry comparable to the operational point of use slurry.

1. Introduction

Chemical mechanical polishing (CMP) is a process that is used to planarize wafers for microelectronic applications. CMP involves polishing the metallic surface of the wafer with a pad while adding slurry. Polishing slurry is one of the most costly consumables in the CMP process. For this reason, the recycling of CMP slurries has become a new challenge for the microelectronic industry and has been investigated for the past 12 years [1–4].

Slurries usually consist of an abrasive material such as silica, alumina or ceria and an aqueous medium that facilitates the suspension of the abrasive particles [5,6]. In the case of tungsten polishing (W-CMP), the aqueous medium contains a strong oxidizer (hydrogen peroxide) and an iron catalyst such as ferric nitrate [7]. The removal of the tungsten layer is achieved by chemical as well as mechanical action. The oxidizer mixture induces the passivation of the tungsten upper layer. The removal of the oxidized tungsten layer requires two processes: passivation/abrasion and chemical dissolution [8,9]. Moreover, it is known that the dissolution static etch rate has to be low to prevent corrosion and planarization defects [10,11]. Other additives may be included in the chemical composition for a better control of the slurry stability and of the tungsten layer formation and removal kinetics [12–14]. The addition of carboxylic acids as complexing agents or stabilizers has been proposed in order to reduce the kinetics of the iron catalyzed oxidation [12,14,15]. The addition of surfactants also helps to stabilize the polishing slurry and prevents settling, flocculation and decomposition of the silica particles [16]. It improves wafer polishing uniformity [14,17,18].

In this study, an attempt is made to recycle commercially available acidic silica-based slurry used for the polishing of tungsten
Abrasive particle and chemical components are highly diluted during the rinsing steps of the CMP process. However, only a small fraction of the slurry is degraded [19] and it could be regenerated through a concentration step. A few studies exist concerning the regeneration of oxide CMP slurry by filtration [20]. In view of the abrasive particle size, the ultrafiltration (UF) process seems to be one of the most suitable processes for the recycling of silica based slurry [21–23]. In addition to the economic interest of recycling slurry, an environmental benefit arises from the potential reduction of waste water volume by ultrafiltration [23]. The problem was that with this process only the abrasive particles are regenerated but the initial chemical composition of the slurry is not recovered. Oxidizing and stabilizing properties are not maintained in the concentrated used slurry (CUFFS). It is a major drawback, especially in the case of W-CMP slurry, for which the chemical composition plays a major role in the chemical and mechanical polishing. As reviewed by several authors, the chemical composition of W-CMP slurry is a determining factor in obtaining uniform and non-defective polishing [5,24,25]. Moreover, in our case, recycling slurry has to be directly operational and all the settings related to the polishing process have to stay unchanged. First experiments were conducted in static conditions where the chemically adjusted slurries were compared to the original slurry as reference [26]. Results confirm the importance of chemistry: (i) the addition of ferric nitrate ensured the passivation of the tungsten layer, obtaining static removal rate similar to that of the reference slurry, but conducted to rapid decomposition of the oxidizer due to the presence of metal impurities in the recycled slurry [27] (ii) the addition of carboxylic acid ensured the increase of the pot lifetime of the recycle slurry (preventing the rapid degradation of hydrogen peroxide) but increased the static etch rate (related to dissolution). From these results it emerged the need to make perfectly balanced chemical adjustments between catalyst and complexing agent in order to ensure the recovering of the specific chemical properties (balance between dissolution and passivation) and the stability of the original slurry [20].

Thus, we performed different chemical adjustments on concentrated used slurry, mainly taking care on the effect of chemistry in order to prevent corrosion and allow soft oxidation of the metal surface by passivation. Carrying out experiments on site under dynamic conditions, we evaluated the influence of additives on the main CMP parameters: removal rate, within wafer non-uniformity and Defectivity, and also the stability of the slurry. Comparing the results obtained under static and dynamic conditions, we were able to discriminate the effect of chemistry on the mechanical removal of tungsten. Validation of the results through design of experiments methodology helps in the identification of the optimal chemical adjustment that has to be made on the recycled slurry to fulfilled industrial polishing requirements to some extent.

2. Experimental

2.1. Slurry concentration and characterization

The reference slurry was silica (SiO₂)-based slurry from Cabot, Semi-Sperse® W2000 (SSW2000), which is used for standard W-CMP process. It contains silica abrasive particles (<200 nm, 1.8 wt.%) in acidic conditions (pH 2.65 ± 0.05) and hydrogen peroxide (2.70 ± 0.05 wt.%) as oxidizer. The slurry to be recycled is collected after use at the outlet of the CMP process. A diverter valve automatically controlled by the polishing time was adapted to the Chemical–Mechanical Planarization system and allowed an effluent to be recovered which was more concentrated than the used effluent [28]. The slurry was retreated in a semi industrial ultrafiltration pilot plant equipped with a module containing hollow polysulphone fibers (S = 1 m², MWCO = 100 kDa, Lp0 = 200 - Lh = 1 m² bar⁻¹) [23]. The filtration flux was stabilized around 50 L h⁻¹ m⁻² for a low Trans Membrane Pressure (TMP = 0.3 bar) implying low energy consumption and fouling. Post filtration was carried out on a polypropylene cartridge (0.5 µm cut-off, Pentair Industrial) in order to eliminate agglomerates. The slurry suspensions – collected used slurry (named US), concentrated by ultrafiltration after used slurry (named CUS) and post-filtrated concentrated slurry (named CUFFS) – were characterized at each step of the recycling process and compared to the point of use (named POU) slurry taken as reference. The slurry characterization was obtained from measurements of physicochemical parameters such as pH (Microprocessor pH-meter HI 221, HANNA Instruments France), conductivity (CDM 210 Conductivity meter with temperature compensation, MeterLab and CDC 745-9 conductivity cells, 2-pole, Radiometer Analytical) and turbidity (Turb 550 IR, WTW).

Dry weight suspensions were obtained after complete dehydration (30 h at 110 °C) of samples in a compact muffle furnace (LE 2/11/ R6, Nabertherm GmbH). The dry solid content was related to the silica concentration only, since the concentration of chemical compounds was low. Turbidity (NTU) was used as an indicator of the amount of suspended solids in the slurry sample. Turbidity values were correlated to silica percentage (wt.%) and gave a good indication of the amount of silica particles in the slurry [22]. Iron content was evaluated by colorimetric o-Phenanthroline method. Hydrogen peroxide titration was carried out using potassium permanganate (ChemLab NV) in acidic solution on a Titroline easy dispenser (SCHOTT Instrument GmbH). The hydrogen peroxide decomposition rate (k) gave an indication of the pot lifetime of the slurry (stability).

Range values for the slurry physicochemical parameters before and after the filtration steps are summarized in Table 1. The concentration of colloidal silica by ultrafiltration led to the regeneration of the physical properties – silica content and particle size distribution – so that they were identical to those of the POU reference slurry. Turbidity and granulometry were analyzed to validate the retreatment process.

The dilution of the slurry during the CMP process also led to the reduction of the conductivity and to an increase in pH. In accordance with previous studies, the silica particles were retained thanks to an ultrafiltration membrane cutoff of 100 kDa whereas the ion concentrations remained constant throughout the filtration step [22,26]. Moreover, the chemical components were only partially recovered due to loss during the CMP cleaning steps and during the ultrafiltration. pH and conductivity were corrected by chemical adjustments. Addition of ferric nitrate, carboxylic acid or H₂O₂ alone tended to decrease the pH and increase the conductivity.

Particle size distributions in volume and in number were estimated by laser granulometry measurements over the particle diameter range 0.6 nm–6 µm on Zetasizer Nano-S (Malvern Instruments). Average particle sizes for the original slurry and the treated slurry, both without dilution, are shown in Table 2.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Physico-chemical characteristics of the point of use slurry and after used and after concentration by ultrafiltration slurry [26].</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physico-chemical parameters</td>
<td>Turbidity (NTU)</td>
</tr>
<tr>
<td>POU</td>
<td>880–990</td>
</tr>
<tr>
<td>US</td>
<td>130–570</td>
</tr>
<tr>
<td>CUS</td>
<td>960–1030</td>
</tr>
<tr>
<td>CUFFS</td>
<td>960–1040</td>
</tr>
</tbody>
</table>
was used and the results were exploited applying response surface analysis using commercial software (NEMROD-W 2000-D LPRAI Corporation, Marseille, France) in order to confirm the influence found elsewhere [26]. According to the previous work, static reference values, 2.70 ± 0.05 wt.% and 2.65 ± 0.05, respectively. These last parameters are pre-requisite for all experiments.

2.2. Chemical adjustments and experimental design

Blanket silica wafers were provided from an industrial user. A thin (8000 Å) tungsten layer was deposited uniformly on each 8 in. wafer by chemical vapor deposition (CVD) at the manufacture. The treated slurry was adjusted with mixed chemicals: (i) ferric nitrate nonahydrate (CAS no. 7782-61-8, Fischer Scientific, UK) used as iron catalyst, (ii) oxalic acid (oxalic acid dihydrate CAS no. 6153-56-6, ACROS Organics) used as iron sequestering agent, (iii) sodium dodecyl sulphate (SDS, acquired from VWR-Prolabo) and Tween® 80 (from Sigma–Aldrich) used as surfactant, and (iv) hydrogen peroxide (30% stock solution from VWR-Prolabo) as oxidizer. Little information could be obtained on the chemical composition of the slurry at the point of use. The reference value for the iron catalyst (Fe) concentration in the slurry was about 0.36 mM. This value was chosen as reference to express all other chemical concentrations in terms of equivalency (1 eq. = 0.36 mM). The hydrogen peroxide concentration and the pH were adjusted as reference values, 2.70 ± 0.05 wt.% and 2.65 ± 0.05, respectively. These last parameters are pre-requisite for all experiments.

Design of experiments (DOE) is particularly useful in our case where uncertainties over the exact composition of the slurry are important. They already proof their efficiency to study CMP process, e.g. they were successfully applied to study the effect of mechanical factors on the removal rate and non-uniformity [29]. Thus, we performed design of experiments (DOE) and variance analysis using commercial software (NEMROD-W 2000-D LPRAI Corporation, Marseille, France) in order to confirm the influence of chemical additives on CMP parameters. A Box-Behnken model was used and the results were exploited applying response surface methodology. Experiment was duplicated at center point so as to provide an indication of inherent variability. Description of variables for DOE is provided in Table 3.

2.3. Chemical mechanical polishing (CMP)

2.3.1. Batch experiments

In static conditions, one single wafer was used for each test in batch experiment. More details of these experiments could be found elsewhere [26]. According to the previous work, static removal rate (SRR) referred as the dissolution of the tungsten layer (static etch rate). SRR is calculated in units of Å min⁻¹ on the basis of the material loss (∆w), the tungsten layer density (ρ) and the wafer surface (A), according to the following equation:

\[
\text{SRR} = \frac{\Delta w}{A \rho t}
\]

2.3.2. Dynamic experiments

A CMP polisher is composed of a base, a turntable covered with a pad (polishing tissue) and a head drive. The slurry particles are conveyed in a regulated flux to the pad where they are retained. The wafer is held in a rack (Fig. 1). The rack and the turntable oscillate. During polishing, the force applied to the rack is transmitted to the wafer which comes into contact with the pad soaked in slurry. The combination of these chemical and mechanical actions allows the planarization of the wafer.

The polishing sequence includes: (i) conditioning of the pad, (ii) polishing, (iii) a rinsing phase and (iv) brushing of the pad. The time and frequency of each phase depend on the type of the polishing process but also on the process recipe chosen.

The pad conditioning consists in removing the top layer of the pad, polluted by the polishing. The pad conditioner is an element in rotation similar to the polishing head and containing of an array of diamonds embedded on nickel plated or steel discs. The mechanical action of the diamond conditioner regenerates asperities and re-opens the pores on the pad surface, improving pad–wafer contact and slurry transport [30].

A 776 Chemical–Mechanical Planarization system with a fully automated Wafer Handling System (WHS) was used as the CMP polisher. The polishing time was set as 1 minute and the polishing sequence was the same as in a production situation. Chemically adjusted slurries were sent onto the pad by the production pump and with a flow around 70 ml min⁻¹. However, there was a difference with the production conditions: the slurry was not pressurized before being sent onto the pad and was not filtered online. The CMP output parameters are generally the removal rate (RR), the within wafer Non Uniformity (NU) and the Defectivity (D). As says were performed at least twice on the same polishing head so as to achieve the best reproducibility possible and to obtain an average value of RR and NU. An average value of Defectivity was obtained using two polished silicon wafers. These parameters were measured on the industrial production equipment (Unpatterned Surface Inspection System, Surfscan SP1-TBI, KLA-Tencor). NU is obtained from the measurement of the difference between the smallest and the greatest thickness all over the wafer and expressed as a percentage compared to the mean thickness. Defectivity characterizes the number of defects occurring on the wafer. Defects are analyzed through an optical method for analyzing light scattering. As defects are often associated to particles present on wafer surface, the sum of all defects is expressed as particle counts.

3. Results and discussion

3.1. Comparison between the reference POU slurry and the recycled CUS slurry

Wafers were polished on site for different chemical mixtures added to the recycled slurry (CUS). The removal rate (RR), the within wafer Non Uniformity (NU), the Defectivity (D) and the hydrogen peroxide decomposition rate (K, h⁻¹) were measured on site. First of all, a comparison was made between the concentrated used slurry (CUS) with only H₂O₂ typical contents addition (implied that H₂O₂ contents is settled at 2.70 ± 0.05 wt.%) and the reference POU slurry. Results are provided in the Fig. 2 (industrial requirements are represented by a shaded area).

Considering the requirements, CUS fulfills the industrial specifications for the main CMP parameters, excepting for Defectivity criteria. No specification is provided for H₂O₂ decomposition rate but a good stability of the slurry is expected. Thus, a K value under 0.01 h⁻¹ or 10% weight loss in H₂O₂ contents within the first hour is the minimum required. CUS, when adding H₂O₂, is less stable than POU with a mean K value about 0.012 ± 0.005 h⁻¹. Even if this

Table 2
Average particle size (nm) in the four slurries studied: POU (n = 14), US (n = 6), CUS (n = 15) and CUS (n = 11) regarding intensity, number and volume distribution.

<table>
<thead>
<tr>
<th>Granulometry data</th>
<th>2.5 Å/Å</th>
<th>Intensity</th>
<th>Number</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>POU</td>
<td>195 ± 25</td>
<td>249 ± 75</td>
<td>135 ± 32</td>
<td>264 ± 35</td>
</tr>
<tr>
<td>US</td>
<td>191 ± 38</td>
<td>232 ± 53</td>
<td>100 ± 48</td>
<td>270 ± 84</td>
</tr>
<tr>
<td>CUS</td>
<td>217 ± 50</td>
<td>343 ± 99</td>
<td>121 ± 58</td>
<td>378 ± 98</td>
</tr>
<tr>
<td>CUSFS</td>
<td>190 ± 34</td>
<td>321 ± 71</td>
<td>103 ± 33</td>
<td>333 ± 94</td>
</tr>
</tbody>
</table>

Table 3
Description of the design of experiments (DOE).

<table>
<thead>
<tr>
<th>Experiment design</th>
<th>Factors (X)</th>
<th>Domains of variation (eq.)</th>
<th>Number of experiments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Box-Behnken</td>
<td>X1: Iron concentration</td>
<td>0–1</td>
<td>13 ± 1</td>
</tr>
<tr>
<td></td>
<td>X2: Oxalic acid concentration</td>
<td>0–2.32</td>
<td></td>
</tr>
<tr>
<td></td>
<td>X3: Surfactant mixture concentration SDS/Tween® 80 (5:1)</td>
<td>0–0.48</td>
<td></td>
</tr>
</tbody>
</table>
The parameter is commonly variable for CUFS or POU as well (more than 40% variability when considering repeated experiments), the difference between recycled (CUFS) and new slurry (POU) is significant. Presence of metal impurities from previous polishing, revealed by the brownish-yellow color of the used slurry, could be responsible for alteration of the slurry stability and modification of the balance between static etch rate and removal rate [27]. Moreover, it could explain the formation of defects related to the precipitation of tungsten oxide impurities [31]. Some little differences can be observed regarding the reference values for all CMP parameters. The mean removal rate (RR) for CUFS is found below the lower value of reference (RR_{CUFS} = 3210 ± 160 Å min⁻¹ against 3560 ± 130 Å min⁻¹ for reference) confirming that the recovering of particles size and concentration using ultrafiltration alone is not enough to obtain efficient removal rate. This point is corroborated by the inverse trend observed for within wafer no-uniformity (NU) and Defectivity (D0). NU values for CUFS are slightly higher than reference (5.1 ± 0.5% and 4.1 ± 1.1%, respectively). As well, Defectivity (D0) is much higher for CUFS, with a mean value of about 690 ± 440 particles, against 160 ± 150 particles for POU. This parameter is of major importance at this point of the process since it will affect the first interconnect metallization layer [31]. Though, the high variability observed for this parameter (more than 60%) makes analysis biased and it could be difficult to assign change in D0 to change in slurry’s composition or intrinsic variation. Thus, Defectivity obtained for CUFS cannot be considered acceptable but further results should be cautiously evaluated. Putting apart this parameter and regarding the high similarity of particles profiles between CUFS and POU slurry (Table 2), all these differences should mainly be attributed to difference in the chemical composition and confirm the need of processing the CUFS with chemical adjustments. The removal rate and the pot lifetime of the recycled slurry have to be increased and the within wafer no-uniformity and the Defectivity have to be decreased to recover the required CMP parameters.

3.2. Effect of chemical additives on CMP parameters

In our previous work, chemical additions were performed on CUFS and experiments were conducted under static conditions. Then, the measured removal rate (static removal rate SRR) is related to the dissolution rate of the tungsten surface layer. Design of experiments DOE was computed to evaluate the influence of chemical additive on the SRR and the pot lifetime (low H₂O₂ decomposition) of the recycled slurry. It was shown that carboxylic acid addition (especially addition of oxalic acid) leads to an increase of the pot lifetime of CUFS. At the contrary, the addition of ferric nitrate rapidly destabilizes the slurry. As well, oxalic acid...
and ferric nitrate had an opposite effect on the SRR, promoting respectively the dissolution or the growth of the tungsten film [26]. It was concluded that chemical adjustments have to be properly balanced in order to insure passivation of the tungsten layer and achieve proper removal rate while slurry stability is under control.

3.2.1. Removal rates and stability of chemically adjusted CUFS

In the frame of design of experiments (DOE) and independent assays, we processed chemical adjustments on CUFS according to different oxalic acid to iron ratios under industrial operating conditions. Chemical adjustments of CUFS are expected to increase the RR and stability with decreasing the NU and DO. The measuring removal rates (RR) are shown in the Fig. 3. Values obtained for the same adjustments in static experiments, as static removal rate (SRR), were added for further analysis of the chemical effect. Error bars were calculated from duplicate experiments and were corrected when conditions were repeated for confirmation purpose.

The dynamic removal rates (RR) are significantly higher than the static removal rate (SRR) as the result of the combinatory effect of chemicals on the dissolution, growth and mechanical removal of the tungsten film. Small addition of iron alone has no observable effect on the global removal rate despite its clear impact on the SRR. Indeed, the dissolution rate decreases when ferric nitrate is added without any effect on the removal rate. At the opposite when no iron is added, the RR decreased out of specification as the oxalic acid addition increased, while a slight increase of the SRR could be observed in comparison to the initial value for CUFS. This means that dissolution is not the principal mechanism in the polishing process but it affects the RR. The higher the dissolution rate, the lower the dynamic removal rate is. Dissolution is observed when H$_2$O$_2$, alone or with oxalic acid, are provided, leading to the formation of a thick and porous oxide tungsten layer [32,33]. Corrosion occurs when dissolution of the passive layer (WO$_3$) into tungstic ions (WO$_4^{2-}$) is initiated, exposing the tungsten layer to further oxidation by H$_2$O$_2$ [34–36]. In these conditions, where dissolution mechanisms are dominant, the mechanical removal is not facilitated and removal rate could not reach the required value. Iron addition, over 0.5 eq., counteracts the oxalic acid addition, diminishing the SRR while the global removal rate increases to the reference POU value. This confirms that the chemical composition of the slurry influences the dynamics of passive film formation, as suggested by Singh and Bajaj [37]. Thus, mechanical polishing efficiency depends on the formation and growth mechanisms of the oxide passive layer (WO$_3$) which are governed by the ferric ions. Albeit, this means that iron is not responsible for dissolution as proposed by Seo et al., it still is critical for the formation of a passivation layer which prevents the tungsten surface from dissolution and corrosion [38]. Without the formation of the passive protective film, there is a risk of default formation during the removal phase by the mechanical actions of particles [39]. Only the mechanical abrasion could remove the soft passivation layer and exposed the tungsten to oxidation process via Fenton’s mechanism as described by Kaufman [40]. Joint reactions of growth and mechanical removal of the passivation layer have a synergistic effect on the global removal rate and should be predominant to provide high removal rate as required [30,41]. Lowest dissolution rate and required global removal rate (3600 ± 40 Å min$^{-1}$) are obtained for an oxalic acid to iron ratio of about 3. These conditions are also optimal for the stability of the slurry, lowering the H$_2$O$_2$ decomposition rate ($K$) under the 0.01 h$^{-1}$ threshold [26]. It confirms that the stabilization of the H$_2$O$_2$ decomposition could be provided by complexing ferric ions by organics such as oxalic acid as proposed by Lim and coworkers [7].

3.2.2. Within wafer non-uniformity and Defectivity of chemically adjusted CUFS

Non-uniformity and Defectivity were measured for the different chemical conditions and results are provided in the Figs. 4 and 5, respectively. Values obtained for POU reference were added for comparison purpose. Within wafer non-uniformity values is found in the specification for most of the chemical conditions.

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**Fig. 3.** Dynamic versus static removal rate (RR versus SRR) for POU and CUFS with different chemical adjustments expressed as the ratio of oxalic acid to iron (OA/I) equivalents addition.
be attributed to a problem with this set of wafers during CVD or Defectivity (experiment without any explanation. This abnormal level of Defectivity. Especially, iron addition most of the tested conditions lead to an acceptable level of Defectivity (>700 part.) for CUFS.

Defectivity value (3.5 ± 0.2%, lower than the reference value. As describe earlier, the acid to iron ratio equal 3), NU is found at an acceptable level shinning. In optimal conditions highlighted in Section 3.2.1 (oxalic acid is added alone to CUFS. The highest NU value is observed when 3 eq. oxalic acid is added. When iron is added, NU is stabilized under the specified threshold value regardless the concentration of oxalic acid. Thus, high NU value could be associated to process drift [27] or, more probably, the generation of local dissolution spots. Indeed, an excess of oxalic acid could be responsible of local passivity break down that could also explain an increase of defects in these conditions [42]. Moreover, it confirms the rule of iron addition to enhance the planarization efficiency. Working against dissolution of the tungsten, iron promotes the uniform passivation of the upper tungsten layer which could be equally and easily removed by the mechanical polishing. In optimal conditions highlighted in Section 3.2.1 (oxalic acid to iron ratio equal 3), NU is found at an acceptable level 3.5 ± 0.2%, lower than the reference value. As describe earlier, Defectivity value (D0) is highly variable and is the only parameter found out of specification (>700 part.) for CUFS.

At the exception of experiments that were conducted at 0.5 eq. iron addition most of the tested conditions lead to an acceptable level of Defectivity. Especially, D0 was higher in the condition of 0.5 eq. iron and 2.3 eq. oxalic acid additions for the duplicate experiment without any explanation. This abnormal level of Defectivity (D0 > 3000 part.) was never observed again and could be attributed to a problem with this set of wafers during CVD or a dysfunction of the polishing process for this set of experiments (probably during the cleaning step when precipitation of metal oxide could occurred due to an increase of pH) [27]. Albeit, no particular trends could be highlight for this parameter, a 0.5 eq. iron should be avoided. Then, oxalic acid and iron adjustments should be done with a ratio equal to 3:1 eq., for which D0 is lowered by 60% (273 ± 40 particles), compared to CUFS without adjustment. In these conditions, D0 is in the industrial specification but is still higher than the reference value. Nowadays, tolerance level for defect is zero.

3.3. Effect of a mixture of surfactants on CMP parameters

Some question remains about how to reduce the Defectivity. Basim and coworkers [43] showed that addition of surfactants, within or near the critical micelle concentration, increases the stability of the slurry and decreases the material removal through the reduction of the friction forces. It was shown that the reduction of particles agglomeration prevents the formation of defects [44]. Moreover, surfactants are used as dissolution inhibitors of the metal surface in chemical mechanical polishing [5,44]. In the case of tungsten CMP, this kind of inhibitor is not particularly required because of the low corrosive potential of tungsten [30], but could help to decrease the excess of dissolution mechanism for tungsten slurry recycling [46,47].

During static experiments, no effect of the surfactants was observed on the H2O2 decomposition rate or the static RR. Nevertheless, some changes were observed on the silica particles dispersion and settling. Mostly, we observed high dispersion and low settling for a mixture of SDS and Tween80 with a ratio of 3:1 for a sum of surfactants inferior to 1 eq. We expected that increasing the dispersion of particles in the slurry by addition of surfactant would improve the within wafer non-uniformity and Defectivity while polishing. Thus, a part of the DOE was conducted to assess the effect of surfactant addition on the CMP parameters. Assays were performed with and without addition of a mixture of surfactants (sum equal to 0.24 eq. or 0.48 eq. SDS:Tween80, 5:1) for CUFS and different oxalic acid to iron ratio conditions. The influence of surfactant addition in CUFS and in the case of optimal oxalic acid to iron ratio conditions (3:1eq.) on CMP parameters (RR, NU and D0) could be observed in the Fig. 6.

Comparison of CMP parameters obtained using CUFS, with and without addition of surfactants, reveals the effect of surfactant additions. It leads to a decrease of the non-uniformity and Defectivity while the removal rate is diminished. This result confirms that these components allow the reduction of the friction forces [18,42]. Whereas Defectivity is found at the highest level when adding oxalic acid alone due to higher dissolution rate, the lowest value of Defectivity (150 ± 15 particles, not shown) was obtained when 0.24 eq. surfactants were added to the worst case conditions of 2.32 eq. oxalic acid in CUFS. Thus, the presence of surfactants counteracts the dissolution effect as an etching inhibitor [45] or diminishes the formation of agglomerates that could be responsible for defects during polishing as suggest by previous studies [48]. At this point a correlation between the antagonist effect of oxalic acid and surfactants addition may be suggested.

In the case of the optimal oxalic acid to iron ratio conditions (3:1 eq.), the addition of surfactants reduces the RR but its effect on non-uniformity and Defectivity is not that significant. No supplemental benefit is obtained by the dissolution inhibitory property of surfactants.

3.4. Validation of the results by the DOE methodology

Part of the results presented above was conducted in the frame of a DOE to assess the statistical significance of the chemical effect
on the CMP parameters RR, NU and D0. Results from duplicate center point experiment are used to provide a measure of inherent variability which is confronted to the variability observed when changing input parameters (chemical additions). The regression in dynamic conditions is less faithful than the static experiments. Even if, there is still some approximation in this exploitation, the response surface methodology provides interesting information from isoresponse curves that confirms some of the previous highlights. Thus, only the most significant results are presented below.

The effect of chemical additions on the RR and NU could be confirmed by DOE. NU is clearly not affected by the chemical adjustments. A significant positive effect is provided by the iron addition alone, whereas surfactants tend to decrease the RR. Oxalic acid added alone has no effect, while combined with an appropriate amount of iron, they significantly increase the RR. Isoresponse curve presented in Fig. 7 shows the synergistic effect of both addition of oxalic acid and iron.

It should be expected that higher concentration, while maintaining the same optimal ratio oxalic acid to iron of 3, would lead to higher removal rate. The statistical analysis for D0 is made difficult because of the high variability of the results obtained for this parameter and validation of the influence of chemical additions on D0 was not possible. Nevertheless, the antagonist effect of surfactants and oxalic acid on Defectivity is clearly shown on the isoresponse curve of D0 in Fig. 8.

Results suggest that surfactants should be added proportionally with oxalic acid in order to preserve low Defectivity. The high variability of D0 values could be explained by the inherent variability and complexity of the polishing process. Thus, results are not surprising since even with POU slurry, the many variations of the output parameters remain unexplained. Besides, it is known that the temperature (which was not monitored during our tests) as well as the applied pressures and rotation speeds (which must be kept constant), are parameters that controls RR [19,27]. Nevertheless, experiments conducted under dynamic condition were carried out in thermostated room. DOE using Box-Behnken model also allowed the determination of the optimal adjustment in order to obtain the best values possible for the three parameters. Table 4 presents the coordinates of this optimal adjustment
(iron: 0.7 eq., oxalic acid: 2.32 eq. and surfactants: 0.47 eq.). Table 5 gives the modeled values for CMP parameters RR, NU and $D_0$ under dynamic conditions and for $H_2O_2$ decomposition rate ($K$).

The overall desirability of this modeling is only 74%. However, the desirability for the Defectivity – parameter which is the most difficult to obtain – is 100%. Besides, it can be noted that this adjustment is very close to the optimal oxalic acid to iron ratio conditions (3:1 eq.). This allows the validity of this optimal adjustment to be confirmed since the response values for the modeled adjustment and the optimal experimental adjustment are very close to one another. Small differences can be attributed to the modeling errors. RR is found in the industrial specifications at the lower range of the reference value. NU is slightly increased but it is not really representative since the input factors studied have no real influence on it. When comparing the slurry with optimal adjustment with CUFS, $D_0$ is slightly decreased by the optimal modeling: the minimal Defectivity obtained with CUFS slurry is close to 600 whereas it is about 150 with the optimal adjustment. Moreover, contrary to CUFS slurry, the slurry with optimal adjustment allows the $H_2O_2$ decomposition rate to decrease and lie within the range obtained for the POU slurry. Thus, the slurry concentrated by ultrafiltration and then chemically adjusted in iron, oxalic acid and surfactant allows maintaining the three main CMP parameters within industrial specifications and is highly stable through time.

Even though these conditions of chemical adjustment almost ensures success, it should be advisable to verify experimentally whether these input factor values are equal to the modeled responses for the CMP output parameters. In order to validate this strategy of reused for tungsten polishing, it is advised to (i) study the Defectivity more specifically with the help of defect mapping and (ii) perform the same type of tests on wafers with chips. Defects review will make it possible to determine the presence of scratches or micro-scratches while the tests on wafers with chips will ensure that the recycling process is in adequacy with further usage in CMP processes.

### 4. Conclusion

In view of reducing the costs of the CMP process and especially of the CMP slurry, the retreatment of slurry is being studied. After the retreatment of the CMP slurry by ultrafiltration to concentrate the silica phase to the same concentration as that of the reference slurry, chemical adjustments are performed to comply with industrial specifications. Thus, we study the effect of three key chemical components: iron, oxalic acid and a surfactant mixture conducting a design of experiments methodology under static and dynamic conditions. These tests are of dual interest. First it is possible to determine the effect of the input factors – chemical additions – on the three main CMP output parameters – dynamic removal rate, non-uniformity and Defectivity. It is also possible to compare these parameters to those obtained with a slurry concentrated without adjustment (CUFS) and to the industrial specifications (POU). Secondly, comparing static and dynamic removal rates, it was possible to distinguish the dissolution of tungsten surface from the global removal rate. This allowed a better understanding of the CMP mechanism regarding the relative influence of chemicals on polishing parameters. Indeed, utilization of static tests could be used as a complement of measures of the corrosion potential [49] and zeta potential [50,51] in the determination of the dissolution/passivation states of the tungsten layer.
Results showed that chemical additions lead to the stabilization of the oxidation process, insure the passivation of the W surface and prevent the formation of defects due to corrosion effects. These influences are highly significant on the removal rate, little significant on the Defectivity and negligible on the within wafer non-uniformity. RR increases with a combined increase in ferric nitrate and oxalic acid concentrations and decreases with surfactant addition. As they decrease RR, the surfactants appear to act as modifiers of the surface state of the wafer and should be able to improve this surface state. An optimization is nevertheless necessary, regarding either the type of surfactant or the type of surfactant mix (anionic or cationic/non-ionic). Whereas, DOE confirms the influence of chemical addition on CMP parameters, it appears that many other variables have an effect on the studied responses during dynamic tests, either other slurry components or parameters of the polishing process.

The DOE produces an optimized scenario in order to adjust the slurry chemistry with good results. CMP parameters obtained in these optimized conditions fulfilled the industrial specifications to some extent. Therefore, it should be considered to recycle the slurry after use by recovering the abrasive particles using ultrafiltration process and by adjusting the chemistry of the slurry (mainly adding the chemical which are lost during the CMP process and the ultrafiltration process).

Finally, in view of reusing slurry in an industrial situation, the polished wafers will be studied and electrical tests performed on the chips. Large-scale testing of a great number of wafers would then allow the validation of the industrial recycling process. This work is focused on specific feature for tungsten metal polishing but regarding the literature \[23\], the proposed approach could be applicable to other material–slurry combinations. Moreover, the capacity to recycle polishing slurries is economically and environmentally beneficial from the reduction of the cost for consumables and the limitation of industrial water rejection and lower solid wastes.

Acknowledgments
This work was supported by the MERISIER project. Also we thank M. Sergent for supporting information on design of experiments and for providing the NEMROD W Software.

References