

New Photopatterning Materials for Advanced Lithography

Jordan Phillip Howard-Jennings, 2018 CNF REU Intern

Engineering, Harvey Mudd College

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CNF REU Principal Investigator: Christopher Kemper Ober, Materials Science and Engineering, Cornell University

CNF REU Mentors: Kazunori Sakai, Materials Science and Engineering, Cornell University; Seok-Heon Jung, Materials Science and Engineering, Cornell University; Christopher Alpha, CNF, Cornell University

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Contact: jhowardjennings@g.hmc.edu, ks2288@cornell.edu, sj736@cornell.edu, alpha@cnf.cornell.edu

Website: http://www.cnf.cornell.edu/cnf_2018reu.html

Abstract:

Photolithography, which is a patterning method used to produce micro/nano-scale features, is currently approaching capabilities of producing sub-10 nm features at EUV wavelengths [1]. We present the findings of material development and characterization of model chemically amplified resists (CARs) to study novel photoacid generators (PAGs), as well as of Zn and Zr metal oxide nanoparticle (MO-NP) resists which offer many advantages over CARs.

Background:

For the past decade, the semiconductor industry has been using immersion lithography to make deep-UV lithography critical dimensions smaller [2]. However, and particularly with CARs which utilize PAGs to “amplify” the photo-reaction that occurs upon exposure, two issues arise: defective patterns resulting from the leaching of resist into the immersion fluid [2], and a sensitivity boost at the expense of line-edge roughness (LER) [3,4]. For EUV lithography to become an industry standard, current materials development of EUV resists must address these issues.

MO-NPs have garnered attention for resist development because they offer many potential and realized advantages to current resist technologies: smaller size relative to polymer photoresists [5], and a ligand-exchange photoreactive mechanism that is more controllable than the deprotection reactions that occur with generated photoacids in conventional resists [6], among other benefits. Currently, metal oxide methacrylate resists produced from hafnium and zirconium have been shown to be potential candidates for EUV resists because of their high resolution and sensitivity [6].

Furthermore, the ligand-exchange mechanism occurring between the acid groups on the metal oxides and the generated photoacids that produces a solubility switch in the exposed area is of interest of further investigation. Since there is a direct correlation between scumming (a phenomena associated with high LER) and the

type of PAG used [3], a study of how PAGs alter resist performance is necessary for understanding how newly developed PAGs may provide better compatibility for EUV MO-NP resists.

Materials and Methods; Synthesis

MO-NP Resists. Resist solutions were prepared by combining 91 mg metal oxide, 9.1 mg PAG (N-hydroxynaphthalimide triflate), and 1.9 g PGMEA in a small glass vial. The metal oxide and PAG were dissolved in PGMEA through 12-16 minutes of stirring on a handheld vortex mixer.

Model CARs. Resist solutions were prepared by synthesizing the monomers tert-butyl methacrylate, isobornyl methacrylate, and methacrylic acid with AIBN as an inhibitor. Once this product was formed, it was mixed with a PAG and PGMEA and dissolved by exposure to a heat gun and vigorous hand-mixing for 10-15 minutes.

Materials and Methods; Lithographic Conditions

MO-NP Resists. Resists were spun onto bare silicon wafers at 2000 rpm for 60s and given a 60s soft bake at 40°C (Zr) or 70°C (Zn). The wafers were then exposed using a 248 KrF source ASML DUV stepper at a dose of

