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Scripting approach in hybrid organic-inorganic condensation simulation: the GPTMS proof-of-concept

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Scripting Approach in Hybrid Organic-Inorganic Condensation Simulation: the GPTMS Proof-of-Concept

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Abstract Silica-based hybrid organic-inorganic materials prepared by sol-gel chemistry exhibit unique chemical and physical properties. (3-glycidoxypropyl)trimethoxysilane (GPTMS)-based networks represent an archetype of this class of substances, with a vast range of applications. In the present study, a new computational recipe has been developed within Materials Studio software platform to generate atomistic models of GPTMS crosslinked networks. The methodology is based on molecular mechanics/dynamics schemes and assumes close proximity as a criterion for crosslinking reaction to occur. The COMPASS force-field was selected for molecular model constructions, and two charge schemes – one obtained directly from the force field and one derived from quantum-chemical calculations – were employed and compared for the prediction of the final system thermophysical properties. Starting from fully-hydrolyzed GPTMS molecules, a realistic three-dimensional network was successfully constructed, including the presence of 4- and 6-membered cyclic structures. Mechanical moduli and specific heat values estimated from equilibrated structures were selected as benchmarks for model/procedure validation. Overall, the simulation results are reasonable and in the range of experimental data available in the literature on similar systems. Thus, the proposed computational strategy has a good potential in the design and optimization of organic-inorganic hybrid materials.

Keywords: condensation reaction, GPTMS, molecular dynamics, Perl scripting, thermophysical properties

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1. Introduction

Nanoscience and nanotechnology are opening new avenues in the fields of chemistry and physics of matter. In particular, the chance to create new, *smart* substances starting from a molecular level clearly constitutes an appealing way to design materials which possess targeted and well-defined macroscopic properties. In this scenario, the molecular building block – a nano-object – is usually an isolated entity (e.g., a molecule or an ensemble of nanoparticles) prepared at the nanometric scale which exhibits a specific chemical and/or

1
2
3 physical property. Sol-gel formation technique constitutes one of the most actual, promising and convenient
4 way to prepare nanomaterials, as it can be produced by using one-step processes which are flexible, efficient,
5 and can be carried out in a wide range of operative conditions. Moreover, different devices characterized by
6 different shapes – matrices, fibers, etc. – can be easily obtained.

7
8 Compared to other chemical elements, silicon is one of the most convenient and productive element for the
9 preparation of the organically modified alkoxides required for the design and synthesis of hybrid organic-
10 inorganic (O/I) materials. A plethora of reasons speak in favor of the peculiar character of silicon, among
11 which the transparency, thermal, and chemical stability of Si-O-Si networks, and the “sweet chemistry”
12 involved in their synthesis (1,2) are just a few. Generally speaking, the polycondensation reaction of
13 alkoxysilanes results in a variety of structures, ranging from monodisperse silica particles to polymer
14 networks, depending of the reaction conditions involved (3). In addition, copolymerization of alkoxysilanes of
15 different functionalities makes it possible to tailor the ultimate material structure and performance. For
16 example, tetrafunctional alkoxysilanes form densely crosslinked silica structure SiO_2 , trifunctional monomers
17 polymerize to branched polysilsesquioxanes (PSSQOs) of the general formula $\text{RSiO}_{3/2}$, whilst bifunctional
18 alkoxysilanes generally yield linear polymer chains $(\text{R}_2\text{SiO})_n$. Ring formation is also a peculiar feature of
19 alkoxysilanes polymerization reactions; clearly, the presence and amount of cyclic structures exert an
20 influence on the ultimate structure and performances of the O/I hybrids (4-7).

21
22 Properties of the microheterogeneous organic-inorganic hybrids depend, to a good extent, on an interphase
23 interaction determining the morphology. Strong interactions, for instance, leads to a reduction of the size of
24 inorganic domains in the organic medium, and often improve the properties. Therefore, organofunctional
25 trialkoxysilane monomers are used to prepare hybrid polymers and are employed as coupling agents, mainly
26 in coating materials. 3-glycidyoxypropyltrimethoxysilane (GPTMS) is an organofunctional alkoxysilane
27 monomer that can undergo both the sol-gel polymerization of the alkoxy groups and curing of the epoxy
28 functionality to form a hybrid network with covalent bonds between organic and inorganic phases. Routinely,
29 however, polymerization of GPTMS is carried out by a sol-gel process which leads to the formation of
30 PSSQO structures with pendant, unreacted epoxy functionalities that are prone to later, eventual curing (4).
31 O/I hybrid materials based on GPTMS have several important applications, including antiscratch coatings (8),
32 contact lens materials (9), passivation layers for microelectronics (10), multifunctional coatings (11), and
33 optical devices (12-14). GPTMS-based optical waveguides, in particular, are very promising materials
34 because of the possibility to incorporate optically active organic molecules in a matrix that is dense at low
35 temperature and with a high degree of microstructural homogeneity (15,16). Accordingly, these extensive
36 range of applications continue to attract various studies on GPTMS hybrid O/I systems. To obtain optimized
37 formulations and efficient technological processes, however, extensive experimental campaigns must be
38 carried out; further, some sound theories in conjunction to experiments must be developed, in order to gain
39 some fundamental knowledge about the physical/chemical phenomena at the basis of the properties of these
40 materials. On the other hand, on the spur of actual industrial competition, the number of lengthy and costly
41 experiments must be drastically reduced, and the establishment of reliable, accurate theories is urgently
42 needed, to be able to design molecular systems with fine-tuned, targeted properties.

43
44 Computer-based molecular simulations nowadays constitutes a versatile, efficient and reliable tool to achieve
45 these goals. Indeed, these techniques can be of great help in reducing experimental hard work by sorting out
46 useless trials and addressing the synthesis and characterization to more productive efforts. Accordingly, in this
47 work we developed a computational strategy to obtain realistic molecular models of crosslinked polymer
48 networks based on a molecular dynamics (MD) Perl script for the Materials Studio software platform. This
49 methodology has been applied to mimick the formation of 3D hybrid O/I networks based on the condensation
50 reaction of GPTMS under acid conditions. Further MD simulations were then carried out to predict some
51 material properties (e.g., elastic constants and heat capacity) of the network structures thus obtained.
52
53
54
55
56
57
58
59
60

2. Computational details

2.1. Model building and general computational recipe

Since the major aim of this work was to develop a script for simulating the generation of an hybrid O/I system based on GPTMS monomer, we decided to start by choosing the simplest conditions leading to the condensation network. Accordingly, we started by modeling a fully hydrolyzed GPTMS molecule, and mimic its condensation reaction under acid conditions (6), i.e., only the creation of Si-O-Si bonds between the available Si-O-H moieties takes place, leaving the epoxy groups unreacted (see Scheme 1).

For the generation of the final crosslinked system, the following general computational recipe was applied:

Step 1. The molecular model of the hydrolyzed GPTMS molecule was built, and its atoms were typed and charged using the COMPASS forcefield (ff) (17, 18). The molecular geometry was then optimized again using COMPASS ff. In order to test the eventual influence of the partial charge distribution on the physical properties of the final crosslinked system, we also assigned to the previously geometry optimized molecule a partial charge scheme obtained recharged using the quantum semiempirical method AM1-ESP as implemented in the *Vamp* toolbox of *Materials Studio*. At the end of **Step 1**, then, two molecular models of the hydrolyzed GPTMS monomer with the identical shape but different partial charges were obtained and used for further calculations.

Step 2. 100 hydrolyzed GPTMS molecules were packed into a simulation box under periodic boundary conditions using the *Amorphous Cell* builder modulus of *Materials Studio*. The initial density of the liquid mixture was set to 1.57 gcm^{-3} , a value estimated from the average experimental density of the final crosslinked system assuming an ideal case of 100% condensation (1.35 gcm^{-3} , 19). To start the condensation reaction/networking process from a representative initial system, 100 different simulation boxes were independently created for each charge scheme considered (i.e., 200 3D cells were obtained overall). After geometry relaxation of each 3D box, the one with the lowest energy value was selected, - namely structures I - one for each charge scheme, for running the molecular dynamics (MD) simulations. Finally, in order to determine the eventual influence of the choice of the initial structure on the simulation results, two further, independent MD simulations for each charging case on structures characterized by approximately the same energy value of structure I after relaxation - namely structures II, were also conducted and analyzed.

Step 3. The selected 3D boxes containing the initial GPTMS monomer systems were first subjected to a geometry optimization (2000 steps); then simulated annealing procedure was applied (20-25) (8 cycles of 1000 MD steps, temperature range 200K - 500K). 5000 steps of molecular dynamics a room temperature (298 K) were carried out. All simulations were conducted in the canonical (NVT) ensemble. After preliminary trials (results not shown), an integration time step of 0.2 fs finally was selected, giving a total annealing time of 1.6 ps, and an overall MD time of 1 ps.

Step 4. The distances between the reactive atoms (O and H atoms from each OH group linked to the Si atom, see Scheme 1) were measured and ordered in an increasing order. The three closest pairs of reactive atoms whose distances were smaller than the selected reactive cutoff distance (3 \AA) were identified, and between the corresponding Si atoms new Si-O-Si bonds were created. An equivalent number of water molecules were deleted from the system (see Figure 1). The reactive cutoff distance was increased during the networking formation from 3 \AA to 6 \AA . Since the presence of small, strained rings (i.e., with less than 4 Si atoms) in the final system is not found experimentally in hybrid O/I systems based on GPTMS (4,26), the necessary restrictions were implemented in the corresponding script, as described in details in the following paragraph.

Step 5. **Steps 3** (referred to the actual cell) and **4** were repeated until no more pairs of reactive atoms satisfying all criteria were detected in the system.

2.2. Condensation reaction script

The whole network formation procedure was achieved by writing a Perl script using the object library available in *Materials Studio* v. 4.1. In what follows, for the sake of clarity and brevity, we will limit the description to the most important technical parts of the final version of the script. Further details are available upon request. Each condensation simulation starts by loading the initial structure obtained at *Step 2* and its association to a proper variable to be used for accessing all objects present in the structure itself. The corresponding command is:

```
my $doc = $Documents{"name.xsd"};
```

The next action consists in deleting all existing sets in the initial structure, i.e.:

```
my $sets = $doc->UnitCell->Sets;
foreach my $set (@$sets) {
  $set->Delete;
}
```

This is an important issue as, in some of the next steps, some new sets will be created with respect to predefined reactive atoms; these sets, in turn, will be used to create the relevant close contacts and, ultimately, bonds. Accordingly, the presence of other sets has to be avoided.

Since the simulations are performed under 3D periodic conditions, one of the filters **DisplayRange**, **UnitCell** or **AsymmetricUnit** should be applied to access any object in the document. These filters define finite sets of accessible objects in the document. For example, the **DisplayRange** filter allows to access the items which are displayed when the document is viewed in the *Materials Studio* Visualizer, whilst the **UnitCell** filter returns objects which are unique with respect to periodic translation of the lattice. Having tested both the **UnitCell** and **DisplayRange** filters, we selected to use the former. From preliminary tests performed on the considered systems (data not shown) it follows that both filters can be employed for scripting. The results are very similar in terms of physical properties of the final structures, and a criterion for the alternative choice of these filters will be the subject of a more detailed study.

It is useful to create a simple variable which will maintain all the atoms instead using

```
$doc->UnitCell->Atoms;
```

This association is done by this command:

```
my $atoms = $doc->UnitCell->Atoms;
```

Another fundamental step that has to be undertaken is proper labeling of all the important chemical entities which, for the specific system considered here, are the linked Si-O-H atoms. Accordingly, all Silicon atoms were labeled **Si**, the oxygen atoms covalently bonded to it **O**, and the reactive hydrogens **H**. The whole Perl code for this action is listed below:

```
foreach my $atom (@$atoms){
  if($atom->ElementSymbol eq "Si"){
    $atom->Name = "Si";
  }
  foreach my $atom1 (@{$atom->AttachedAtoms}){
```

```

1
2
3         if( $atom1->ElementSymbol eq "O" && $atom1->Name ne "RO"){ $atom1->Name = "O";
4
5     foreach my $atom2 (@{$atom1->AttachedAtoms}){
6         if( $atom2->ElementSymbol eq "H"){ $atom2->Name = "H";}
7
8             }
9
10        }
11
12    }
13
14 }
15

```

The **RO** label was chosen for the oxygen atoms bridging two Si atoms (i.e., Si-O-Si). Since it may be necessary to restart the whole script several times, it is essential to avoid relabeling **RO** oxygens to **O**, and this justifies the choice of the label **RO** in the above piece of code.

Before performing any calculation, all relevant parameters for Geometry Optimization (GO), Annealing (AN) and Molecular Dynamics (MD) must be set, and each proper setting saved to a file (*.xms). This setting file must be placed later in the same directory where the Perl script is located. Then, the command for loading the parameter file:

```

25 Modules->Forcite->LoadSettings("name");
26

```

must be included in the script. This avoids the tedious repeating of setup operations. As can be seen from the command above, the *Forcite* module of *Materials Studio* was employed in the calculations performed in this work, as scripting cannot be implemented in the *Discover* engine available in version 4.1 of *Materials Studio*. Direct commands inside the script are then used to set some important calculation parameters for running the calculations. The settings and running commands for GO, AN, and MD) calculations employed in the present study are reported below:

```

35 Modules->Forcite->ChangeSettings(["ChargeAssignment" => "Use current"]);
36

```

```

37 Modules->Forcite->Calculation->Run($doc, Settings(Task => "Geometry Optimization", MaxIterations => 2000));
38

```

```

39 Modules->Forcite->Calculation->Run($doc, Settings(Task => "Anneal", TimeStep => 0.2, AnnealCycles => 8,
40 EnergyDeviation => 2000000 ));
41

```

```

42 Modules->Forcite->Calculation->Run($doc, Settings(Task => "Dynamics", TimeStep => 0.2, Ensemble3D =>
43 "NVT", NumberOfSteps => 5000, EnergyDeviation => 2000000 ));
44

```

When a trajectory file (*.xtd) is created as output of the AN and MD simulations, the **\$doc** variable is automatically reinitialized, and from that moment it represents the whole trajectory of the system. Since just the last frame from each of these trajectories is needed as input for subsequent calculations, this last frame is used for reinitializing the document variable **\$doc** after each AN/MD run. Furthermore, in the case of MD runs (after which *Step 4* is applied, see above), another important reason to update **\$doc** variable to the last frame of the trajectory can be envisaged. Since some atoms are to be deleted from the system during the condensation reaction leading to network formation, (i.e., H₂O molecules, see Figure 1), the atomistic file (*.xsd) and not the trajectory file (*.xtd) has to be used as, once the number of atoms in actual frame is altered, the entire trajectory is invalidated. The complete sequence of commands in AN, MD part of our script is as follows:

```

56 Modules->Forcite->Calculation->Run($doc, Settings(Task => "Anneal", TimeStep => 0.2, AnnealCycles => 8,
57 EnergyDeviation => 2000000 ));
58
59
60

```

```

1
2
3 my $numFrames = $doc->Trajectory->NumFrames;
4 $doc->Trajectory->CurrentFrame = $numFrames;
5 my $LastFrame = Documents->New("After-ANN.xsd");
6     $LastFrame -> CopyFrom($doc);
7 $doc -> Discard;
8 $doc = $LastFrame;
9
10 Modules->Forcite->Calculation->Run($doc, Settings(Task => "Dynamics", TimeStep => 0.2, Ensemble3D => "NVT",
11 NumberOfSteps => 5000, EnergyDeviation => 2000000 ));
12

```

```

13 my $numFrames = $doc->Trajectory->NumFrames;
14 $doc->Trajectory->CurrentFrame = $numFrames;
15 my $LastFrame = Documents->New("After-MD.xsd");
16     $LastFrame -> CopyFrom($doc);
17 $doc -> Discard;
18 $doc = $LastFrame;
19

```

On the first line of the **\$doc** updating code, the new variable **\$numFrames** is created, and the number of frames of the considered trajectory is assigned to it. Then, the last frame of the trajectory is set as the actual one. Afterwards, the empty atomistic file is created (**After-ANN.xsd** or **After-MD.xsd** in the above routine) and associated to the variable **\$LastFrame**. In the next line of the code, the last frame of the actual trajectory is copied into the new atomistic file; subsequently, the current trajectory variable is discarded, clearing this document (i.e., the trajectory file) from the computer memory. This obviously does not imply that the trajectory file which is represented by this variable is deleted. Each trajectory file, together with the corresponding (**After-ANN.xsd** or **After-MD.xsd**) files, are saved in the corresponding *Materials Studio* project. The last step is updating the document variable **\$doc** by variable **\$LastFrame**, where the last frame from the previous trajectory is stored.

Since *Steps 3* and *4* are repeated several time during the entire simulation recipe, the **After-ANN.xsd** and **After-MD.xsd** files are automatically numbered (e.g., **After-ANN (2)**, **After-ANN (3)** etc.) by *Materials Studio*. The same automatic procedure is applied to the corresponding trajectories files. No saving instructions then need to be implemented since *Materials Studio* includes all created files automatically into the open project.

Now let us describe a little more in details *Step 4*, which includes i) the analysis of the distances of the pairs of reactive atoms, ii) the creation of new Si-O-Si bonds, and iii) the deletion of H₂O molecules from the system. In order to make the system react, the optimal close contacts between the atoms involved in the condensation reaction have to be realized; accordingly, the atom pairs between which close contacts should take place have to be defined. Given the chemistry of the systems considered in this work, where condensation takes place between Si-O-H···H-O-Si groups (see Scheme 1), only O – H close contacts are needed, where O is the name for the oxygens bound to the Si atoms, and H designs the hydrogens linked to the O oxygens. The part of our script that creates these close contacts is written below:

```

47 # Create arrays to store the reactive atoms in
48

```

```

49 my @reactive_O;
50 my @reactive_H;
51
52 foreach my $atom (@$atoms) {
53     if ($atom->Name eq "O" ) {
54         push (@reactive_O, $atom);
55     } elsif ($atom->Name eq "H" ) {
56         push (@reactive_H, $atom);
57     } else {
58         push (@reactive_O, $atom);
59
60

```



```

1
2
3         push (@reactive_H, $atom);
4     }
5 }
6

```

The above piece of code creates two atom arrays, **reactive_O** and **reactive_H**, which, intuitively, store all atoms except atoms named **H** and named **O**, respectively. Since close contact restrictions can be applied only to sets, however, it is necessary to create relevant sets based on previously created atomic arrays. This can be done by the following commands:

```

13 my $reactive_O_set=$doc->CreateSet("reactive_O",\@reactive_O);
14 my $reactive_H_set=$doc->CreateSet("reactive_H",\@reactive_H);
15

```

Then, the **ExclusionMode** is set to 'Set' value, which should guarantee that creation of close contacts between atoms inside each defined set is forbidden. Clearly in this study only the close contacts between atoms named **O** and **H** are expected, since these atoms are in different sets. The command that make the above described exclusion mode setting is the following:

```

21 Tools->BondCalculation->ChangeSettings([ExclusionMode => 'Set']);
22

```

Unfortunately, this elegant trick works perfectly only under non-periodic conditions (e.g., a non-periodic superstructure). In the case of a system under periodic boundary conditions, such as in the present work, it is necessary to introduce some further filtering to eliminate all the non **O – H** close contacts (see the code part for the creation of **GoodCloseContacts** array listed below). Nevertheless, given the 3D symmetry characteristics of our system, this approach can be successfully applied, ultimately resulting in a substantial reduction computer memory.

The last step necessary for calculating close contacts is the setting of **DistanceCriterionMode**, **MinAbsoluteDistance** and **MaxAbsoluteDistance** parameters. In our case, the following setting was employed:

```

34 Tools->BondCalculation->ChangeSettings(Settings( DistanceCriterionMode =>"Absolute",
35 MinAbsoluteDistance => 0.0, MaxAbsoluteDistance => 3));
36

```

At this point, predefined close contacts can be calculated according to:

```

39 $doc->CalculateCloseContacts;
40

```

Again, it is appropriate to create a variable(s) which will represent the set of created close contacts: **\$CloseContacts**:

```

45 my $CloseContacts = $doc->UnitCell ->CloseContacts;
46

```

For the next procedure, mainly in order to save significant time and memory in loop operations, the creation and use of an arrays, such as **GoodCloseContacts**, in which all the necessary information can be stored, needs to be created:

```

51 my @GoodCloseContacts;
52

```

The code for the initialization of the **GoodCloseContacts** array is listed below:

```

55 foreach my $CloseContact (@$CloseContacts) {
56     my $Name1= $CloseContact->Atom1->Name;
57     my $Name2 = $CloseContact->Atom2->Name;
58
59
60

```

```

1
2
3 my $Atom1 = $CloseContact->Atom1;
4 my $Atom2 = $CloseContact->Atom2;
5 if ( ($Name1 eq "O") && ($Name2 eq "H") || ($Name1 eq "H") && ($Name2 eq "O") ) {
6     push(@GoodCloseContacts,[1,$Name1,$Name2,$Atom1->X,$Atom1->Y,$Atom1->Z,$Atom2->X,
7     $Atom2->Y,$Atom2->Z,$CloseContact->Length,$CloseContact->Atom1,$CloseContact->Atom2]);
8     }
9 }
10

```

In the above code, each element of the **GoodCloseContacts** array has the structure [**status**,**Name1**,**Name2**,**x1**,**y1**,**z1**,**x2**,**y2**,**z2**,**Length**,**Atom1**,**Atom2**], where **status** represents a variable which denotes if a given close contact can take place or not at a given instant. **Name1** and **Name2** are the names of the atoms which belong to that given close contact. **x1**, **y1**, **z1**, **x2**, **y2**, **z2** are the coordinates of these atoms, **Length** denotes the length of the given close contact, and finally **Atom1** and **Atom2** are the atom objects which belongs to the close contact. The names of these atoms are stored in the array as separate variables (i.e., **Name1** and **Name2**); however, they are accessible through the atom objects. This last action was taken just to save some time, since atom names will be frequently used in the analysis section (see below).

It is also worthwhile to note that the filtering of non **O–H** close contacts discussed previously is included in the piece of code listed above; accordingly, the **GoodCloseContacts** array contains only **O–H** close contacts. However, in order to simulate a network-forming condensation reaction, only those **O–H** close contacts which are created between two different GPTMS molecules must be taken into account. Therefore, it is necessary to filter out the **O–H** close contacts which are created between O and H atoms belonging to the same GPTMS molecule. This can be simply achieved by analyzing the array **GoodCloseContacts** and checking whether the species **Atom1** and **Atom2** belong to the same GPTMS molecule or not. In the affirmative, the variable **status** must be changed from 1 to –1, and all the subsequent analysis must be carried out only on those close contacts which have variable **status** equal to 1.

The way to identify the molecular object to which a given atom belongs is using:

```

33 $atom1->Ancestors->Molecule
34

```

Unfortunately, at present, there is no direct method to obtain the index of any given molecule in a molecular model. Thus, a way to circumvent the problem is to identify two different molecules by their different center of mass. To access the coordinates of the center of mass of any given molecule, the following construction can be used:

```

41 my $x1 = $atom1->Ancestors->Molecule->Center->X;
42 my $y1 = $atom1->Ancestors->Molecule->Center->Y;
43 my $z1 = $atom1->Ancestors->Molecule->Center->Z;
44

```

Finally, the **GoodCloseContacts** is ordered by increasing close contact length which, in each element of this array, is represented by the **Length** variable.

At this point, the new Si-O-Si bonds building procedure can begin. As the whole code for this section is big, and the detailed description of each part will exceed the scope of this article, it will only be summarized and briefly commented below.

The first close contact (by index) in the array **GoodCloseContacts** which has the **status** variable equal to 1 (i.e., the shortest one, see above) is chosen. Two new variables **\$O** and **\$H** are created, which are initialized by the atoms **Atom1** and **Atom2** from the selected **GoodCloseContacts** element. All the remaining atoms involved in the bond formation are then identified, and assigned to the proper variables (**\$H1**, **\$O1**, **\$Si**, and **\$Si1**, respectively). Figure 2 illustrates the association of these variables to the corresponding atoms.

1
2
3 According to the bonding scheme of Figure 2, at the beginning only the two atoms which belongs to the
4 chosen close contact (**\$O** and **\$H**, respectively) are considered. Next, the hydrogen atom **\$H1**, which will be
5 deleted together with atoms **\$O** and **\$H**, from the system due to the formation of a H₂O molecule, must be
6 identified (see Figure 2). It is important to note that, in the real process, the water formed during the
7 condensation reaction remains in the reaction environment, and eventually contributes to the hydrolysis of
8 nonhydrolyzed GPTMS molecules. In the system considered here, however, the GPTMS monomers were
9 fully hydrolyzed (a realistic condition which can be easily achieved experimentally). Therefore, water plays
10 no additional role in the process, and does not need to be further considered. Next, atoms **\$O1**, **\$Si**, and **\$Si1**
11 must be identified, in order to create the new bond **\$Si-\$O1** (see Figure 2). This identification process can be
12 carried out resorting to the **AttachedAtoms** function. Below, the piece of code for the identification of the **\$H1**
13 and **\$Si** atoms is listed, as an example:
14
15
16

```
17 my $Si;  
18 my $H1;  
19 foreach my $at (@{$O->AttachedAtoms}){  
20 if($at->Name eq "Si"){ $Si=$at;} else { $H1=$at;}  
21 }  
22
```

23
24 As mention in the introduction, although the structure of a GPTMS-based hybrid O/I network obtained under
25 acid catalysis is similar to that of a crosslinked polymer network, some cyclic structures can be present (4,26).
26 In order to apply some restrictions related to ring size in the formation of eventual cyclic structures, a check
27 whether the bonding between the **\$Si** and **\$O1** atoms will not result in the closure of a strained ring (i.e., with
28 less than 4 Si atoms) must be performed. This leads to the necessity of identifying atom **\$Si1** (see Figure 2);
29 this operation could also serve graphical purposes: for example, the need of changing graphical style for both
30 bonds **\$Si-\$O1** and **\$O1-\$Si1** in order to differentiate them, for example, from other bonds.
31 Should every condition described above be satisfied, the new bond **\$Si - \$O1** is created at this stage with the
32 command:
33
34

```
35 $doc->CreateBond($Si, $O1, "Single");  
36
```

37 otherwise, the next available close contact in the **GoodCloseContacts** array is selected, and the whole
38 procedure outlined above is repeated.

39 As said, once the new Si-O bond is created, a water molecule is released. In our model, this corresponds to
40 deleting atoms **\$O**, **\$H**, and **\$H1** from the system. To accomplish this goal, instead of deleting the required
41 atoms directly (e.g., using commands such as **\$O->Delete**, etc., which can generate problems, especially when
42 using periodic boundary conditions, as once some individual atoms are deleted, all its periodic images are also
43 deleted from the set of close contact atoms).
44
45

```
46 push (@AtomsToDelete,$H);  
47 push (@AtomsToDelete,$O);  
48 push (@AtomsToDelete,$H1);  
49
```

50 After the end of the **\$i** loop, a set based on the **AtomToDelete** array is created, and all the atoms which are
51 deleted from the system according to the reaction scheme will be cancelled at once and in a safe way,
52 according to:
53
54

```
55 my $AtomsToDelete_Set=$doc->CreateSet("AtomsToDelete",\@AtomsToDelete);  
56 $AtomsToDelete_Set->Items->Delete;  
57  
58  
59  
60
```

Once the atoms **\$O**, **\$H**, and **\$H1** are added to **AtomsToDelete** array, and the **\$Si - \$O1** bond is created, it is appropriate to analyze the whole **GoodCloseContacts** array from the actual index **\$i** to the end, and switch the variable **status** to -1 in all those cases in which some of the atoms **\$O**, **\$O1**, **\$H**, and **\$H1** are periodically equivalent to **Atom1** or **Atom2**.

For checking periodic equivalence of two atom, the following function was created:

```

10 sub AtomsArePeriodicEquivalent {
11     my ( $A1, $A2) = @_ ;
12     if($A1->Name ne $A2->Name){ return 0;}
13     my @dA = ($A2->X - $A1->X,$A2->Y - $A1->Y,$A2->Z - $A1->Z);
14     my @length = ($lengthX,$lengthY,$lengthZ);
15     for(my $i=0;$i<3;$i++){
16         if(abs($dA[$i])!=0 && abs($dA[$i])!=$length[$i]) {return 0;}
17     }
18     return 1;
19 }
20 #The variables $lengthX, $lengthY and $lengthZ denotes the length of the sides of the periodic box.
21

```

Figure 3 shows that, if the mentioned **GoodCloseContacts** revision is checked only for equality and not for periodic equivalence of relevant atoms, unrealistic valences of oxygen and silicon atoms are generated.

There can be several configurations which originate these problems, and example of which is illustrated in Figure 4. In the left part of Figure 4 it can be seen that, when more reactive centers are in close proximity, we can see some “collision ” possibility if more reactive centers is in proximity. For example, suppose that the O-H contact (i.e., black O and blue H) is shorter than the other O-H contact (i.e, green O and blue H). Under this condition, the bond between the blue O and black Si is created (red line). If no check of the remaining **GoodCloseContacts** elements is performed, then also the Si-O bond between the green Si and blue O is created. If only equality check after creation of the blue O and black Si bond is performed, no problem arises if the configuration corresponds to the portrayed on the left side of Figure 4. However, if the position of the green molecule is shifted to the periodic image position, then both O-H atoms (the green O and the small blue H atom) are not equal to any other atom (the black and large blue H atom, and the black and the blue O atom), but the small blue H atom is periodically equivalent to that large one, which is a sufficient condition for creating the Si-O bond (blue O and green Si atoms) which is interrupted by the periodic cell boundary but can be clearly seen using the InCell viewtype. This situation is nicely illustrated in Figure 5, taken from the real system of Figure 3.

Then, the bonding procedure with respect of the next O – H close contact from array **GoodCloseContacts** which has **status** equal to 1(or, more precisely, the **GoodCloseContacts[\$i][0]** element) must be repeated.

After a desired number of bonds (**\$MaxNumBond**) is reached, or at the end of the **GoodCloseContacts** array is reached, all sets created and populated in the previous steps must be deleted to release computer memory. Also, renaming of the bridging oxygens (**\$O1**) must be performed, since these atoms must not enter the next **\$I** loop for creating new O - H close contacts.

To give a final overview of the entire procedure, the most critical steps are summarized below using a pseudocode:

```

53 ***START OF THE SCRIPT***
54

```

```

55 #Loading of the structure (*.xsd)
56

```

```

57 my $IterMax=50;           #number of the main loops
58
59
60

```

```

1
2
3 my $MaxNumBond=3;           #maximum number of created bonds in one I loop
4
5
6 #Main loop
7 for(my $I=0;$I<$IterMax;$I++){
8
9     Loading of the setup file (*.xms)
10    Geometry Optimisation
11    Annealing
12    Molecular Dynamics
13    Creation of the reactive atom arrays and sets
14    Close contacts calculation
15    Creation of GoodCloseContacts array and close contacts filtering
16    Ordering of the GoodCloseContacts elements by increasing close contact length
17
18        for (my $i=0;(($NumOfCreatedBonds<$MaxNumBond) && ($i<=#GoodCloseContacts));$i++){
19
20
21
22            Identification of all the atoms $O, $H, $H1, $O1, $Si and $Si1
23
24            Creation of $Si-$O1 bond
25            Addition of atoms $O, $H, $H1 to array AtomsToDelete
26            GoodCloseContacts array revision
27
28        } #end of the $I loop (Step 4)
29
30    #Deletion of reacted atom sets
31    $reactive_O_set->Delete;
32    $reactive_H_set->Delete;
33
34    #Deletion of all calculated close contacts
35    $CloseContacts->Delete;
36
37    #Creation of the set $AtomsToDelete_Set from the array AtomsToDelete and appropriate deletion of all the
38    atoms marked in previous steps for deletion
39    #the if statement excludes attempt to create set from the empty array
40
41    if($#AtomsToDelete + 1>=1){
42    my $AtomsToDelete_Set=$doc->CreateSet("AtomsToDelete",@AtomsToDelete);
43    $AtomsToDelete_Set->Items->Delete;
44    }
45
46    #Renaming of oxygen atoms involved in the formation of Si-O-Si bonds (i.e., accessed using the variable $O1)
47    from "O" to "RO"
48
49    foreach my $atom (@$atoms )
50    {
51    if($atom->Name eq "O"){
52        my $NumOfSi=0;
53        foreach my $atom1 (@{$atom->AttachedAtoms})
54        {
55            if($atom1->Name eq "Si"){ $NumOfSi++;}
56        }#end $atom1
57    if($NumOfSi==2){$atom->Name="RO";}
58    }
59
60

```

```

1
2
3 #end of if($atom->Name eq "O"
4 }
5 #end of loop $atom
6 }
7 #end of I loop (main loop, which should be repeated $IterMax times)
8
9
10 ***END OF THE SCRIPT***
11

```

At the very end it may be worth noticing that, despite we release all computer memory properly in each **\$I** loop, during each script run the memory used by the computer is slowly and progressively increasing, so that, after a number of **\$I** loops, the limit of accessible memory is reached and calculations have to be manually restarted. The number of successfully accomplished **\$I** loops clearly depends upon the available RAM of a given computer, but also on the amount of close contacts which are created during each **\$I** loop. That is the reason why we increase **MaxAbsoluteDistance** from 3 to 6 Å.

2.3. 3D network relaxation for thermophysical properties determination

After the GPTMS 3D network structures were built starting from the two partial charge scheme and applying the script detailed above (i.e., two structures I and two structures II), each system was subjected to further annealing cycles up to 600K. The minimum energy structure for each system was selected for further NVT and NPT molecular dynamics simulation at 300K for data collection. The velocity Verlet algorithm was used for integration in all MD simulations. A time step of 0.2 fs, and the Nosè/Berendsen thermostat were employed for NPT and NVT MD simulations, respectively. The cut-off for non-bond interactions was set at 9.50 Å. The final density of the MD equilibrated systems corresponding to the two different atomic partial charge schemes, AM1-ESP and COMPASS, respectively, is reported in Table 1. As can be seen from Table 1, the equilibrated network densities obtained from initial structure modeled with both partial charge schemes are utterly similar; further, starting from two different annealed initial GTPMS 3D boxes (i.e., structure I and structure II), leads to the same network final density values, thus validating the initial structure selection protocol.

2.4. Mechanical properties (elastic constants) determination

The mechanical behavior of a given molecular system can be described by using continuum mechanics. Since, however, any molecular system has a discrete structure, the model to be employed for the estimation of the elastic constants is an equivalent-continuum model (27), in which the overall mechanical response of representative volume elements to an applied set of boundary conditions is equivalent to the response of the molecular representative volume system subjected to the same set of boundary conditions. The equivalent-continuum is assumed to have a linear-elastic constitutive behavior. The generalized constitutive equation of the equivalent continuum hence is given by:

$$\sigma_{ij} = C_{ijkl} \varepsilon_{kl} \quad (1)$$

where σ_{ij} are the components of the stress tensor ($i, j = 1, 2, 3$), C_{ijkl} are the components of the linear-elastic stiffness tensor, and ε_{kl} are the components of the strain tensor. It is further assumed that the system has isotropic material symmetry.

In atomistic calculation, the internal stress tensor in a system can be obtained using the virial expression:

$$\sigma = -\frac{1}{V_0} \left[\left(\sum_{i=1}^N m_i (\mathbf{v}_i \mathbf{v}_i^T) \right) + \left(\sum_{i<j} r_{ij} \mathbf{f}_{ij}^T \right) \right] \quad (2)$$

where index i runs over all particles through N , m_i and \mathbf{v}_i are the mass and the velocity of the particle, respectively, \mathbf{f}_i is the force acting on the particle, and V_0 is the undeformed system volume.

Basically, two classes of methods for calculating material elastic constants using molecular simulations are available in literature at the present. Consistently, the static method (i.e., based on molecular mechanics) was found to be more practical and reliable than the one based on molecular dynamics (28). Thus, a constant strain minimization method, belonging to the class of static methods, was applied to the equilibrated 3D O/I network system. Accordingly, after an initial system energy minimization, three tensile and three pure shear small deformations (to remain within elastic limits) are applied. The system is then again energy minimized following each deformation. The stiffness matrix is calculated from the second derivative of the potential energy U with respect to strain ε as follows:

$$C_{ij} = \frac{1}{V} \frac{\partial^2 U}{\partial \varepsilon_i \partial \varepsilon_j} = \frac{\partial \sigma_i}{\partial \varepsilon_j} = \frac{\sigma_{i+} - \sigma_{i-}}{2 \varepsilon_j} \quad (3)$$

where σ_i is the i th component of the internal stress tensor, and σ_{i+} and σ_{i-} are the components associated with the stress tensor under tension and compression, respectively (further details on the static method are available in (29)). The Lamé constants for the structure - λ and μ - can in turn be calculated from the related stiffness matrix:

$$\lambda = \frac{1}{3} (C_{11} + C_{22} + C_{33}) - \frac{2}{3} (C_{44} + C_{55} + C_{66}) \quad (4)$$

$$\mu = \frac{1}{3} (C_{44} + C_{55} + C_{66}) \quad (5)$$

For isotropic materials, the stress-strain behavior can be finally be described in terms of the Lamé constants according to the equations:

$$E = \mu \frac{3\lambda + 2\mu}{\lambda + \mu} \quad (6)$$

$$G = \mu \quad (7)$$

$$B = \lambda + \frac{2}{3} \mu \quad (8)$$

$$\nu = \frac{\lambda}{2(\lambda + \mu)} \quad (9)$$

where E , G , B , and ν represent Young's modulus, shear modulus, bulk modulus and Poisson's ratio, respectively.

2.5. Constant pressure and constant volume heat capacities determination

Statistical fluctuations about the mean values of quantities measured during the course of an MD simulation can be directly related to thermodynamic properties. Beside common average quantities like density, pressure,

or energy, the analysis of fluctuations allows to determine properties like heat capacities, compressibility, thermal expansion coefficient, or the Joule-Thomson coefficient. Specific heat capacity at constant pressure C_p , for instance, is obtained from the fluctuations of energy in the isothermal-isobaric ensemble (NPT) according to:

$$C_p = \frac{1}{kT^2} \langle \partial(K + U + pV)^2 \rangle \quad (10)$$

where k is the Boltzmann constant, K and U denote the instantaneous values of the kinetic and potential energy, respectively, and T , P , and V are the familiar thermodynamic state variables. In addition, the notation δX stands for $X - \langle X \rangle$, where $\langle X \rangle$ denotes the ensemble average value of a given quantity X .

3. Results and discussion

3.1. Structural analysis

Figure 6 shows the geometry optimized molecular model of the GPTMS molecule, along with the atom labeling used in this work. For this monomer, the relevant structural parameters are listed in Table 2. These include the Si-O and Si-C1 bond lengths, and the O-Si-O and O-Si-C bond angles, which were estimated to be 1.670 Å, 1.909 Å, 108.4°, and 109.5°, respectively. Unfortunately, experimental X-ray diffraction or other spectroscopy data for GPTMS are not available to date. Therefore, to further validate the geometrical features of our model we applied the same model building/optimization procedure to structurally related molecules, for which such information could be retrieved from the literature. The relevant names, structural formulas and geometrical parameters are listed in Table 3.

By comparing the numbers in Tables 2 and 3, given the differences in molecular species and the fact that, whilst ff calculations are performed on an isolated molecule *in vacuum* the corresponding experimental quantities are obtained either from single-crystal or spectroscopic studies, we can conclude that all our results are in good agreement with the literature data.

In order to check the influence of the atomic partial charges on the final properties of the system, the GPTMS optimized structure was also assigned a partial charge distribution as derived from AM1-ESP calculations (see Table 4). Generally, speaking, a common procedure to estimate atomic partial charges is via Mulliken analysis (33). However, since the results may be strongly dependent on the basis set employed, and there is no unambiguous method to assign charge to two atoms within a given bond (34), we tried to bypass the problem by fitting the point charges at preselected positions to the electrostatic potential surface (ESP) (35,36). On the other hand, the partial charge scheme thus obtained may be, in turn, dependent on the specific molecular conformation considered (37). To verify whether this was our case, we tested the obtained charge distribution by calculating molecular dipole moments and standard enthalpies of formation for all molecules listed in Table 3, and compared them with the corresponding experimental values, where available (see Table 5). As results from this Table, once again the agreement between calculate and experimental quantities is good, thus confirming the validity of the adopted approach.

According to the procedure outline above, an hybrid O/I 3D system based on GPTMS was successfully generated with high conversion. It is well known that a 100% conversion is rarely achieved experimentally because of gel transition at later stage. Although other systems with a different (lower) conversion degree α could be generated by changing, for instance, the distance between close contacts up to a reasonable value of 10 Å, the amount of unreacted group in the actual simulated molecular systems is, on average, equal to 10%, yielding $\alpha = 0.9$.

1
2
3 The central unit cells of the initial GPTMS monomers, and the final network systems obtained from both
4 structures I and II are shown in Figures 4, 5 and 6, respectively. As can be seen from Figures 5 and 6, the 3D
5 network structures is characterized by the presence of both chemical and physical crosslinks. Some bonds
6 clearly connect to image cells across the boundary, and thus extend throughout the periodic system. Due to the
7 scarcity of experimental characterization, which by the way accounts for the small number of simulation
8 studies on crosslinked polymeric systems (39-41), only a few comparisons between experimental and
9 simulation can be attempted based on the main structural features of the systems.

10 As mentioned in the introduction, PSSQOs are the generic products obtained by the hydrolytic condensation
11 of monomers such as GPTMS. Strictly speaking, the term refers to fully condensed structures of formula
12 $(\text{RSiO}_{3/2})_n$ ($n = \text{even number}$), also denoted T_n . But the term is frequently extended to denote partially
13 condensed structures, $[\text{RSiO}_{3/2-x}(\text{OH})_{2x}]_m$ or $T_n(\text{OH})_m$, where $m = 0, 2, 4 \dots (2+n)$ for $n = \text{even number}$ and $m =$
14 $1, 3, 5 \dots (2+n)$ for $n = \text{odd number}$. Keeping this definition, PSSQO structures may vary from perfect
15 polyhedra, incompletely condensed polyhedral, ladder polymers, open structures, linear polymers, and all
16 possible combinations thereof. Generally speaking, then, according to the polymerization conditions
17 employed different structures for the resulting networks are proposed, such as randomly connected three-
18 dimensional networks of trifunctional monomers, "ladder" structures, and a combination of linear, "ladder"
19 and cage-like fragments (42).

20 As the presence and relative amount of the structures described above significantly affect the network final
21 structure, homogeneity and, ultimately, the mechanical properties of the resulting hybrids, the different
22 reaction steps can be optimized to obtain final materials with targeted properties. Experimentally, it has been
23 found that, under acid conditions, only a very small amount of cage-like structures are formed in the GPTMS
24 polymerization, the intermolecular condensation being the preferred mechanism of network growth (6).
25 Accordingly, a high-molecular weight branched PSSQO grows, until a gel system is formed at a conversion
26 degree approximately equal to 0.7. The corresponding composition of the gel state exhibits a considerable
27 amount of silicon atoms involved in triple intermolecularly branched units representing the cross-links in the
28 network. Further SAXS experiments revealed also that, under acid catalysis, no microphase separation
29 resulting in self-assembly of regularly arranged domains is present (6). Contrarily, a broad distribution of
30 high-molecular weight PSSQOs with dangling organic substituents is formed, the structure not promoting any
31 ordering. The inspection of the 3D structures obtained from our simulation procedure compare well with the
32 network picture described above. Indeed, the overall network structure is linear and extends in all three
33 directions, the presence of unstrained rings with a number of Si atoms greater than 4 are present. To further
34 confirm the quality of the network structures, Table 6 lists the mean values of some geometrical parameters,
35 as obtained from initial structure I with the COMPASS atomic partial charges scheme by averaging over 10
36 MD frames. By comparing these data with the corresponding geometrical features of the GPTMS monomer
37 model reported in Table 2, we can see that the differences are rather small, indeed confirming that
38 crosslinking does not result in considerable geometrical modifications, and no substantial strain is induced in
39 the final network.

40 41 42 43 44 45 46 47 48 49 **3.2. Thermophysical properties of the hybrid O/I network**

50 51 52 **3.2.1. Mechanical properties (elastic constants)**

53 The mechanical properties calculated using the 3D network structures obtained from the MD simulations are
54 reported in Table 7. The second column refers to the properties of the network structures simulated with the
55 partial charge scheme obtained using the AM1-ESP approach, whilst the third column lists the same quantities
56 resulting from the 3D structures bearing the partial charges as assigned by the COMPASS ff (see Table 4). In
57
58
59
60

1
2
3 the last column of this Table, the range of experimental values of the corresponding mechanical properties
4 (43,44) is also reported for comparison.

5
6 The data in Table 7 indicate that the charging method has a certain impact on the mechanical properties of the
7 final material. Indeed, in the case of the AM1-ESP partial charging scheme the values obtained from the
8 simulation are located close to the lower limit of the expected range, if not below. With the COMPASS force
9 field charge scheme, the mechanical characteristic of the GPTMS network lay in the expected range.
10 Although no experimental measures are available for our systems, as a conclusion we could say that the
11 predictions obtained by using a standard force field satisfactorily reproduce the available mechanical
12 experimental data.
13

14 15 16 3.2.2. Heat capacities

17
18 The specific heat capacities at constant volume and constant pressure, c_v and c_p , calculated for the GPTMS-
19 based O/I network are reported in Table 8. Again, these properties were obtained for structure bearing the two
20 different partial charge schemes (see second and third column of Table 8 for AM1-ESP and COMPASS
21 charge scheme, respectively). In the last column, the experimental data range available in literature for similar
22 systems is also shown for comparison (43,44).
23

24 As seen for the mechanical properties, all data calculated starting from structures bearing atomic partial
25 charges derived from the AM1-ESP approach lay in the lower limit of the experimental data range, whilst
26 those obtained from the COMPASS partial charge set fall better within the interval of observed values.
27
28
29
30

31 4. Conclusions

32
33 In this work a computational procedure, based on a Perl script developed in Materials Studio v. 4.1, is
34 presented, which allowed to obtain atomistic models of hybrid organic/inorganic networks based on GPTMS.
35 The actual procedure, however, can be applied with minor modifications to any similar system. Furthermore,
36 the translation of the script in other programming languages (e.g., C++, Fortran, etc.) makes this script platform
37 independent and, coupled to an MD routine, allows the same scientific results to be achieved with other
38 simulation codes. The simulations allowed also to estimate several thermophysical properties of the 3D
39 network, such as elastic constants and specific heats, whose values were found to be in the range of the
40 corresponding experimental evidence for similar systems. These results confirm both the accuracy of the 3D
41 model structure generated by the script procedure, and the quality of the force field used. Therefore, the
42 proposed computational recipe can constitute a useful tool for the design and development of new hybrid O/I
43 systems with specific structural/chemico-physical properties.
44
45
46
47

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References

1. Sanchez, C.; Soler-Illia, G.J.d.A.A.; Ribot, F.; Lalot, T.; Mayer, C.R.; Cabuil, V. *Chem. Mater.* 2001, 13, 3061.
2. Livage, J.; Sanchez, C. *J. Non-Cryst. Solids* 1992, 145, 11.
3. Kelts, L.W.; Armstrong, N.J. *J. Mater. Res.* 1989, 4, 423.
4. Matejka, L.; Dukh, O.; Brus, J.; Simonsick, W.J.Jr.; Meissner, B. *J. Non-Cryst. Solids* 2000, 270, 34.
5. Eisenber, P.; Erra-Balsells, R.; Ishikawa, Y.; Lucas, J.C.; Mauri, A.N.; Nonami, H.; Riccardi, C.C.; Williams, R.J.J. *Macromolecules* 2000, 33, 1940.
6. Matejka, L.; Dukh, O.; Hlavata, D.; Meissner, B.; Brus, J. *Macromolecules* 2001, 34, 6904.
7. Brus, J.; Spirkova, M.; Hlavata, D.; Strachota, A. *Macromolecules* 2004, 37, 1346.
8. Nass, R.; Arpac, E.; Glaubitt, W.; Schmidt, H. *J. Non-Cryst. Solids* 1990, 121, 370.
9. Philipp, G.; Schmidt, H. *J. Non-Cryst. Solids* 1984, 63, 283.
10. Popall, M.; Kappel, J.; Pilz, M.; Schulz, J.; Feyder, G. *J. Sol-Gel Sci. Technol.* 1994, 2, 157.
11. Schmidt, H. *J. Non-Cryst. Solids* 1994, 178, 302.
12. Sorek, Y.; Zevin, M.; Reisfeld, R.; Hurvits, T.; Rushin, S. *Chem. Mater.* 1997, 9, 670.
13. Knobbe, E.T.; Dunn, B.; Fuqua, P.D.; Nishida, F. *Appl. Opt.* 1990, 29, 2729.
14. Sorek, Y.; Reisfeld, R.; Tenne, R. *Chem. Phys. Lett.* 1994, 227, 235.
15. (a) Guglielmi, M.; Brusatin, G.; Della Giustina G. *J. Non-Cryst. Solids* 2007, 353, 1681. (b) Della Giustina, G.; Brusatin, G.; Guglielmi M.; Romanato F. *Mat. Sci. Eng. C* 2007, 27, 1382. (c) Brusatin, G.; Della Giustina, G.; Guglielmi, M.; Casalboni, M.; Proposito, P.; Schutzmann S.; Roma G. *Mat. Sci. Eng. C* 2007, 27, 1022.
16. Zevin, M.; Reisfeld, R. *Opt. Mater.* 1997, 8, 37.
17. Sun, H.; Rigby, D. *Spectrochim. Acta Part A* 1997, 53, 1301.
18. Sun, H. *Macromolecules* 1995, 28, 701.
19. Brusatin, G., personal communication.
20. Scocchi, G.; Posocco, P.; Fermeglia, M.; Pricl, S. *J. Phys. Chem. B.* 2007, 111, 2143.
21. Fermeglia, M.; Cosoli, M.; Ferrone, M.; Piccarolo, S.; Mensitieri, G.; Pricl, S. *Polymer* 2006, 47, 5979.
22. Fermeglia, M.; Ferrone, M.; Pricl, S. *Mol. Simulation* 2004, 30, 289.
23. Toth, R.; Coslanich, A.; Ferrone, M.; Fermeglia, M.; Pricl, S.; Miertus, S.; Chiellini, E. *Polymer* 2004, 45, 8075.
24. Fermeglia, M.; Ferrone, M.; Pricl, S. *Fluid Phase Equilib.* 2003, 212, 315.
25. Fermeglia, M.; Pricl, S. *AIChE J.* 1999, 45, 2619.
26. Fidalgo, A.; Ilharco, L.M. *J. Non-Cryst. Solids* 2001, 283, 144.
27. Odegard, G.M.; Gates, T.S.; Nicholson, L.M.; Wise, K.E. *Compos. Sci. Technol.* 2002, 62, 1869.
28. Raaska, T.; Niemela, J.S.; Sundholm, F. *Macromolecules* 1994, 27, 5751.
29. Theodorou, D.N.; Suter, U.W. *Macromolecules* 1986, 191, 139.
30. *Handbook of Chemistry and Physics*, Lide D.R. Ed., 79th ed., CRC Press, Boca Raton, FL, USA, 1999
31. Barrow, M.J.; Ebsworth, E.A.; Harding, M.M. *Acta Crystal.* 1979, B35, 2091.
32. Koput, J.; Wierzbicki, A. *J. Mol. Spectrosc.* 1983, 99, 116.
33. Mulliken, R.S. *J. Chem. Phys.* 1955, 23, 1833.
34. Singh, U.C.; Kollman, P.A. *J. Comput. Chem.* 1984, 5, 129.
35. Weiner, S.J.; Kolmann, P.A.; Nguyen, D.T.; Case, D.A. *J. Comput. Chem.* 1986, 7, 230.

- 1
- 2
- 3
- 4 36. Weiner, S.J.; Kolmann, P.A.; Case, D.A.; Singh, U.C.; Ghio, C.; Alagona, G., Profeta, S.; Weiner, P.J. J.
- 5 Am. Chem. Soc. 1984, 106, 765.
- 6 37. Reynolds, C.A.; Essex, J.W.; Richards, W.J. J. Am. Chem. Soc. 1992, 114, 9075.
- 7 38. Girard-Reydet, E.; Riccardi, C.C.; Sautereau, H.; Pascault, J.P. Macromolecules 1995, 28, 7599.
- 8 39. Fan, H.B.; Yuen, M.M.F. Polymer 2007, 48, 2174.
- 9 40. Wu, C.; Xu, W. Polymer 2006, 47, 6004.
- 10 41. Yarowsky, I.; Evans, E. Polymer 2002, 43, 963.
- 11 42. Boury, B.; Corriu, R. Chem. Rec. 2003, 3, 120.
- 12 43. Van Krevelen, D.W. Properties of polymers, Elsevier, Amsterdam, The Netherlands (1990).
- 13 44. Bicerano, J. Prediction of polymers properties, M. Dekker, New York, USA (1996).
- 14
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Table 1. MD equilibrated density values ρ for the 3D GPTMS-based O/I network structures obtained from the simulations.

Atomic partial charge scheme	Initial structure I ρ (g/cm ³)	Initial structure II ρ (g/cm ³)
AM1-ESP	1.368	1.369
COMPASS	1.371	1.368

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Table 2. Geometrical parameters of the GPTMS molecular model after geometry optimization with COMPASS ff. Atom numbering as in Figure 6.

Si-O1(O2,O3)/Å	Si-C1/Å	H1-O1/Å	C1-C2/Å	C2-C3/Å	C3-O4/Å	O4-C4/Å	C4-C5/Å	C5(C6)-O5/Å	C5-C6/Å	H4-C1/Å	H12-C5/Å
1.670	1.909	1.110	1.531	1.527	1.419	1.419	1.495	1.428	1.439	1.104	1.098
O1-Si-O2	H1-O1-Si	Si-C1-C2	C1-C2-C3	C2-C3-O4	C3-O4-C4	O4-C4-C5	C4-C5-C6	C5-O5-C6	H6-C2-H7	H13-C6-H14	
108.4°	109.5°	114.7°	115.1°	106.6°	114.2°	105.9°	125.4°	60.2°	107.2°	112.2°	

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Table 3. Computed geometrical data for molecules structurally related to GPTMS. Experimental available data (30-32) are reported in parenthesis for comparison.

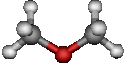

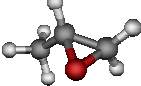
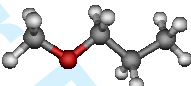
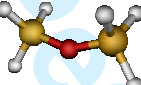
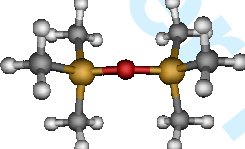
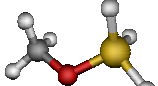
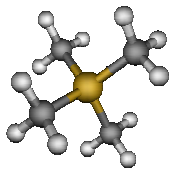
Name	Chemical Formula	Chemical Structure	Geometrical parameters						
Dimethyl ether	C ₂ H ₆ O		C-O/Å 1.412 (1.416)	C-H/Å 1.124° (1.121°)	C-O-C 111.8° (112°)	H-C-C 108.3° (108°)			
Ethylene oxide	C ₂ H ₄ O		C-C/Å 1.459 (1.466)	C-H/Å 1.096 (1.085)	C-O/Å 1.428 (1.431)	C-O-C 60.4°	C-C-H 116.5° (116.6°)		
Propylene oxide	C ₃ H ₆ O		C(H ₃)- C(H)/Å 1.527 (1.510)	C(H)- C(H ₂)/Å 1.452	C-O/Å 1.426	C-O-C 60.4°	C-C-C 122° (121°)		
Propyl Methyl Ether	C ₄ H ₁₀ O		C-O/Å 1.415 (1.418)	C-C/Å 1.527 (1.520)	C-H/Å 1.115 (1.118)	C-O-C 112.3° (111.9°)	O-C-C 109.7° (109.4°)	H-C-H 109.2° (109°)	
Disiloxane	H ₆ OSi ₂		Si-O/Å 1.635 (1.63) ^a	Si-H/Å 1.485 (1.490)	Si-O-Si 151.3° (151.2°) ^b	H-Si-H 109.8°			
Hexamethyl-disiloxane	C ₆ H ₁₈ OSi ₂		Si-O/Å 1.639 (1.638)	Si-C/Å 1.862 (1.869)	H-C/Å 1.101 (1.104)	C-Si-O 108.3° (109.0°)	Si-O-Si 149.7° (151.3°)	C-Si-C 110.8°	
Methoxysilane	CH ₃ OSi		Si-C/Å 1.421	Si-O/Å 1.669	H-C/Å 1.101	H-Si/Å 1.476	C-O-Si 122.6°	H-Si-H 110.8°	
Tetramethylsilane	C ₄ H ₁₂ Si		Si-C/Å 1.880 (1.875)	H-C/Å 1.110 (1.115)	C-Si-C 109.5°	H-C-H 109.7° (109.8°)			

Table 4. Partial charges on GPTMS atoms as assigned by COMPASS and obtained from AM1-ESP calculations. Atom numbering as in Figure 6.

Atom name	Si	O1	O2	O3	O4	O5	C1	C2	C3	C4	C5	C6		
COMPASS	0.80	-0.47	-0.47	-0.47	-0.32	-0.32	-0.24	-0.11	0.054	0.054	0.11	0.054		
AM1-ESP	0.72	-0.27	-0.32	-0.30	-0.10	-0.099	-0.25	-0.20	-0.22	0.14	-0.12	-0.052		
Atom name	H1	H2	H3	H4	H5	H6	H7	H8	H9	H10	H11	H12	H13	H14
COMPASS	0.25	0.25	0.25	0.053	0.053	0.053	0.053	0.053	0.053	0.053	0.053	0.053	0.053	0.053
AM1-ESP	0.13	0.12	0.12	0.045	0.064	0.10	0.093	0.084	0.096	0.016	-0.014	0.070	0.076	0.072

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Table 5. Computed dipole moment and standard enthalpy of formation for molecules structurally related to GPTMS. Experimental available data are reported in parenthesis for comparison.

Name	μ_{calc} (D)	μ_{exp} (D)	$\Delta H_{\text{form,calc}}$ (kcal/mol)	$\Delta H_{\text{form,exp}}$ (kcal/mol)
Dimethyl ether	1.36	1.30	-44.66	-44.00
Ethylene oxide	1.95	1.89	-13.61	-12.57
Propylene oxide	2.06	2.01	-21.68	-22.63
Propyl Methyl Ether	1.13	1.11	-55.08	-56.91
Disiloxane	0.34	0.24	-65.22	–
Hexamethyldisiloxane	0.38	–	-182.01	-185.87
Methoxysilane	1.1	1.15	-53.98	–
Tetramethylsilane	0.00	0.00	-57.99	-57.12

Table 6. Mean geometrical parameters of the GPTMS 3D network obtained from structure I with the COMPASS atomic partial charges scheme by averaging over 10 MD frames.

Si-O/Å	Si-C1/Å	O-Si-O	O-Si-C	Si-C-C	Si-O-Si
1.660	1.928	124.1°	104.3°	123.5°	142.8°

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Table 7. Young modulus E , bulk modulus B , shear modulus G , Poisson ratio ν , and Lamé constants λ and μ for the 3D GPTMS-based O/I network structures obtained from MD simulations.

Mechanical properties	Structure I AM1-ESP	Structure II AM1-ESP	Structure I COMPASS	Structure II COMPASS	Experimental data ^a
E (GPa)	2.85	3.00	3.52	3.96	1.9-4.5
B (GPa)	1.89	2.00	2.58	2.66	2.5-7
G (GPa)	1.14	1.20	1.38	1.58	0.9-1.5
ν	0.25	0.27	0.27	0.25	0.3-0.4
λ (GPa)	1.13	1.20	1.66	1.61	-
μ (GPa)	1.14	1.20	1.38	1.58	-

^aAverage values for crosslinked networks

Table 8. Specific constant pressure (c_p) heat capacity for the 3D GPTMS-based O/I network structures obtained from the MD simulations.

Specific heat capacity	Structure I AM1-ESP	Structure II AM1-ESP	Structure I COMPASS	Structure II COMPASS	Experimental data ^a
c_p (kJ/kgK)	1.16	1.23	1.64	1.49	1.1-2.2

^aAverage values for crosslinked networks

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Figure Captions

Scheme 1. Reaction leading to the Si-O-Si network formation.

Figure 1. Details of the GPTMS molecular modeling showing, in stick-and-ball representation, the reactive atom pairs (top), and the newly formed Si-O-Si bond and the corresponding deleted water molecule (bottom). All other atoms are in line rendering. Color code: Si, gold; O, red; C, gray, H, white.

Figure 2. Atom/variable association in the bonding scheme considered in this work.

Figure 3. GPTMS crosslinked structure created using only equality (not equivalence) check of the atoms during the **GoodCloseContacts** revision. The unrealistic oxygen atoms forming three bonds with Si are highlighted in green, whilst the Si atoms with 4 oxygen bonds are highlighted in violet.

Figure 4. Schema of one problematic configuration leading to unrealistic valences of O and Si atoms (see text for details).

Figure 5. Details of the real system illustrated in Figure 3. Top: **InCell** lattice display style view; bottom: **default** lattice display style view.

Figure 6. Geometry optimized molecular model of the GPTMS molecule, along with the atom labeling used in this work. All atoms are in stick-and-ball representation, using the same color code of Figure 1.

Figure 7. Central unit cell of the initial GPTMS monomers. Color code as in Figure 1.

Figure 8. Hybrid O/I 3D network structures obtained starting from GPTMS models bearing atomic partial charges calculated with the AM1-ESP method (structure I, top left, structure II, top right), and assigned by the COMPASS ff (structure I, bottom left, structure II, bottom right). (see text for details). Crosslinked Si and O atoms are highlighted in Stick-and-Ball. Color code as in Figure 1.

Figure 9. Details of the hybrid O/I 3D network structures obtained starting from GPTMS models bearing atomic partial charges calculated with the AM1-ESP method (structure I, top left, structure II, top right), and assigned by the COMPASS ff (structure I, bottom left, structure II, bottom right). (see text for details). Crosslinked Si and O atoms are highlighted in Stick-and-Ball. Color code as in Figure 1.

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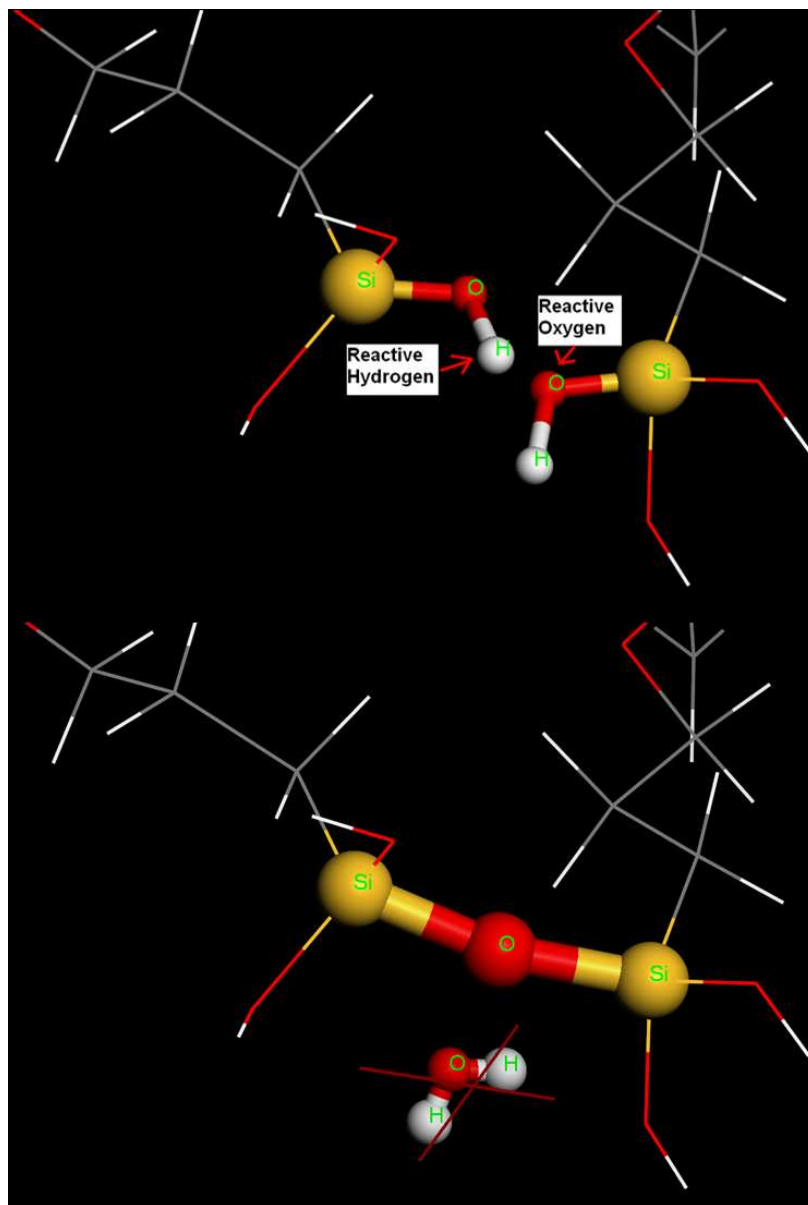
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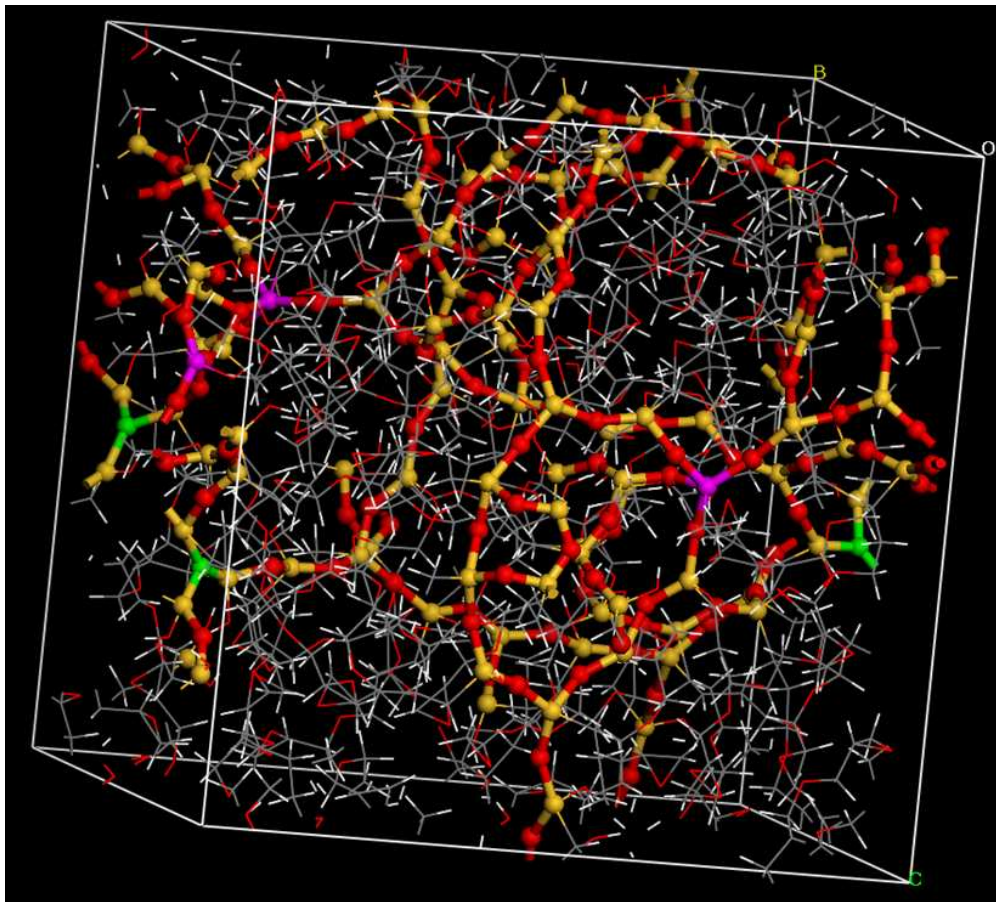
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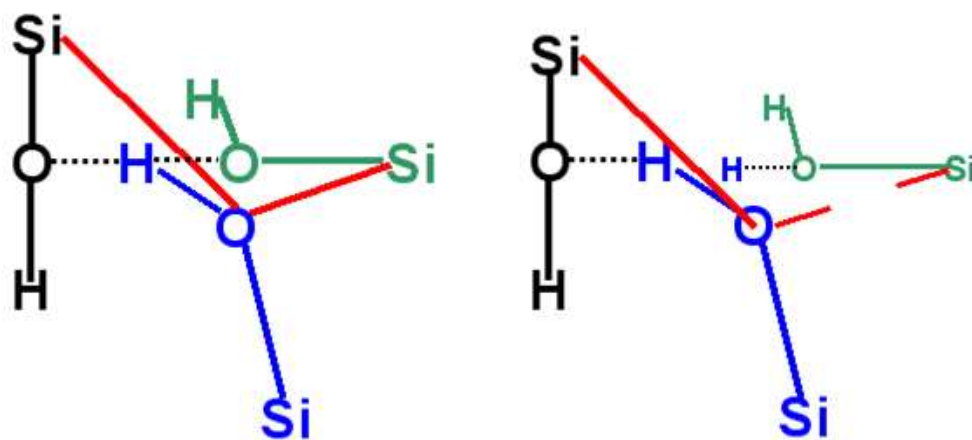
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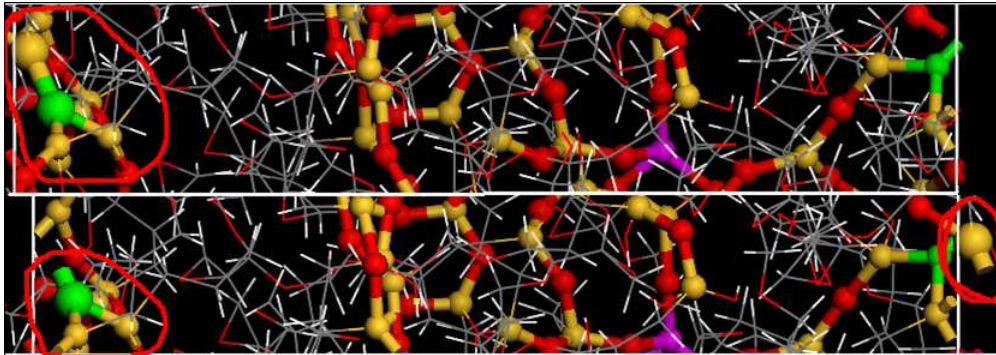
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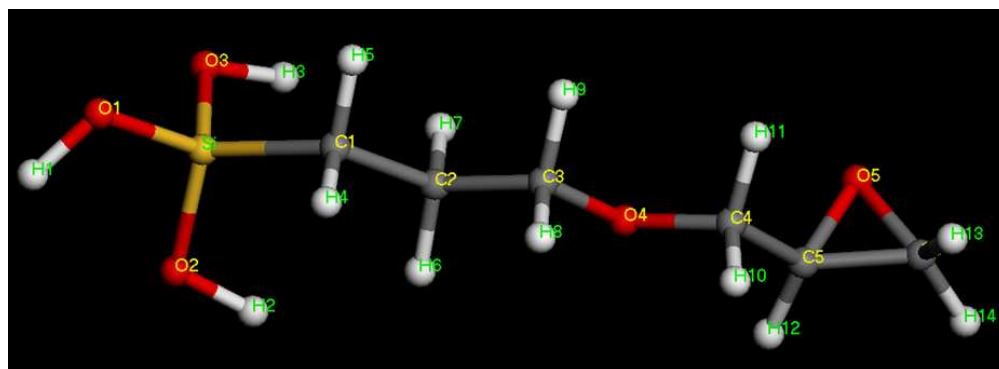
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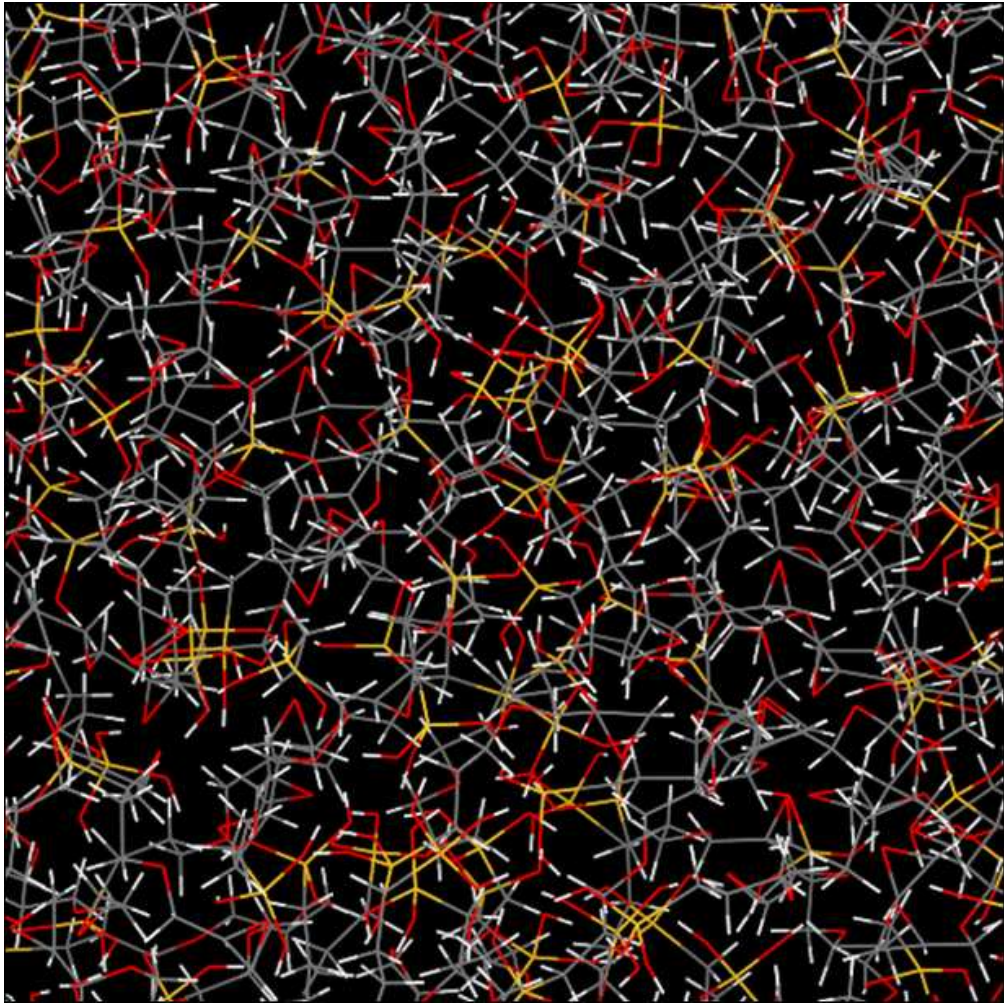
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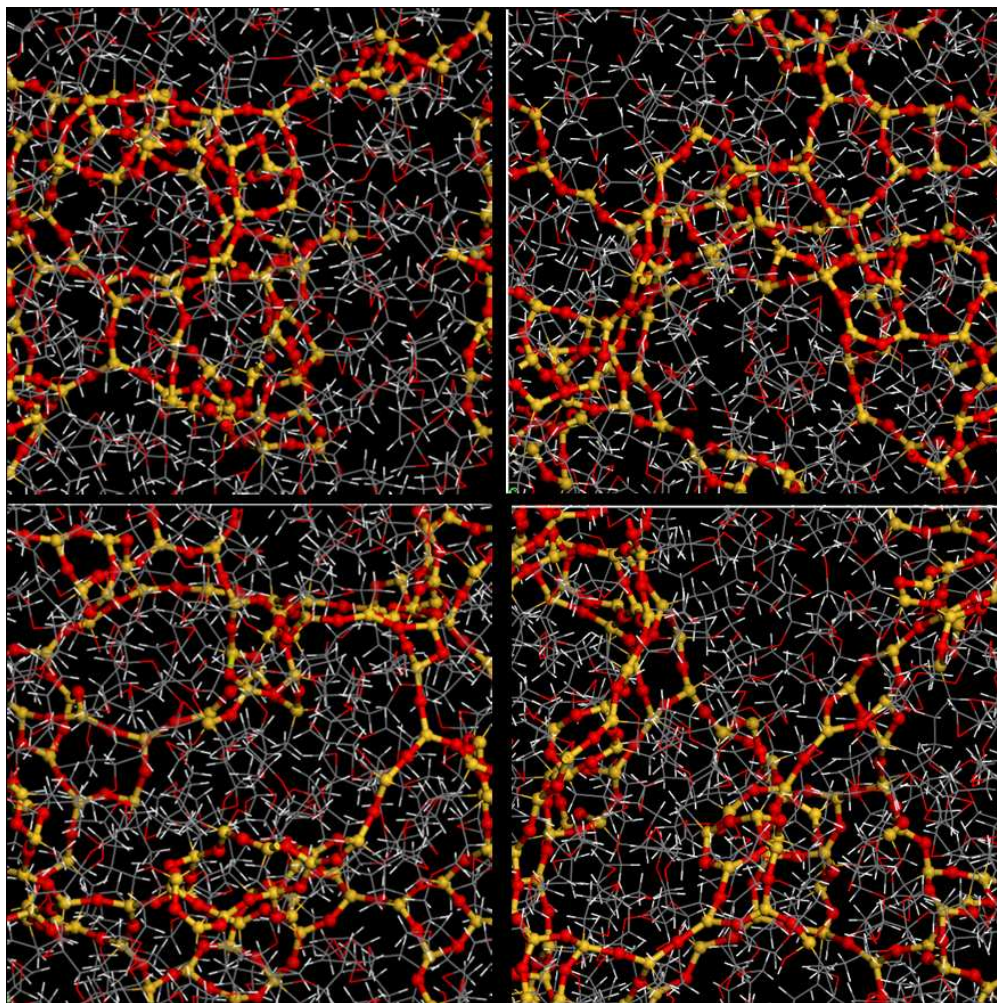
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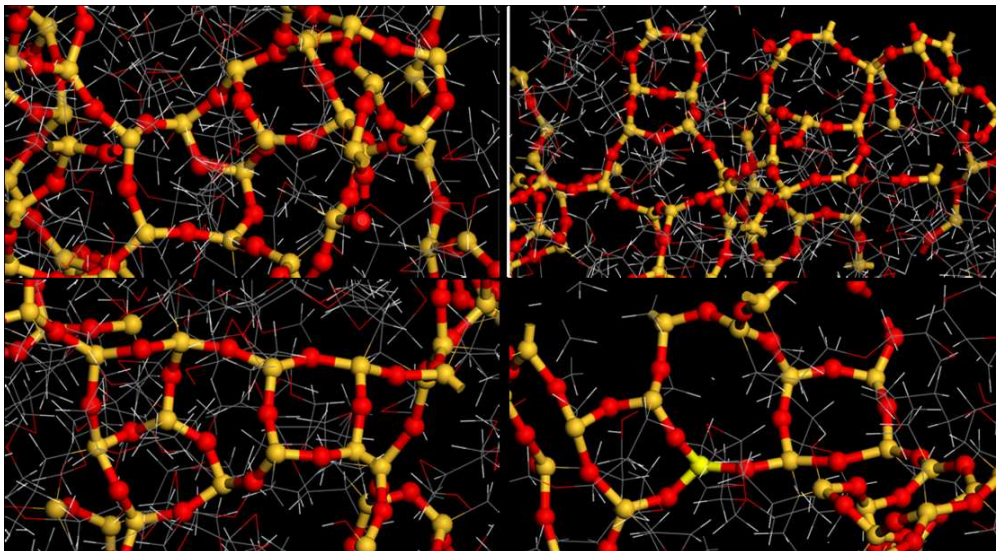
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