Supporting Information:

Coarse-grained modelling to predict the packing of porous organic cages

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1 Parameters for the coarse-grained model

Table S1: Interaction parameters used in the window-to-window simulations.

| $\sigma_{\text{ang}}$ | $J$ (kJmol$^{-1}$) |
|-----------------------|--------------------|
| 0.1                   | 96                 |
| 0.2                   | 56                 |
| 0.3                   | 46                 |
| 0.4                   | 41                 |
| 0.5                   | 36                 |
| 0.6                   | 30                 |
| 0.7                   | 25                 |
| 0.8                   | 21                 |
| 0.9                   | 18                 |
| 1.0                   | 17                 |

Table S2: Interaction parameters used in the window-to-arene simulations.

| $\sigma_{\text{ang}}$ | $J$ (kJmol$^{-1}$) |
|-----------------------|--------------------|
| 0.1                   | -                  |
| 0.2                   | 73                 |
| 0.3                   | 50                 |
| 0.4                   | 38                 |
| 0.5                   | 28                 |
| 0.6                   | 19                 |
| 0.7                   | 15                 |
| 0.8                   | 12                 |
| 0.9                   | 10                 |
| 1.0                   | 9                  |

Table S3: Values of the offset value for the torsional modulation term used in each simulation.

| Packing type         | Offset angle |
|----------------------|--------------|
| window-to-window     | $\pi/2$      |
| window-to-arene      | 0            |
2 Sizes of clusters for phase determination

Table S4: Cluster sizes for phase determination in the window-to-window simulations.

| k_BT | 0.1 | 0.2 | 0.3 | 0.4 | 0.5 | 0.6 | 0.7 | 0.8 | 0.9 | 1.0 |
|------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
|      |     |     |     |     |     |     |     |     |     |     |
| 1.04 | -   | -   | -   | -   | -   | -   | -   | -   | -   | 117 |
| 0.99 | 139 | 391 | 397 | 399 | 354 | 159 | 218 | 100 | 142 | 257 |
| 0.94 | 365 | 407 | 433 | 429 | 412 | 288 | 292 | 247 | 201 | 270 |
| 0.90 | 224 | 433 | 439 | 437 | 417 | 376 | 344 | 300 | 304 | 330 |
| 0.85 | 414 | 435 | 441 | 444 | 439 | 394 | 395 | 383 | 332 | 357 |
| 0.81 | 453 | 442 | 437 | 447 | 419 | 410 | 396 | 406 | 392 | 373 |
| 0.77 | 222 | 438 | 441 | 453 | 415 | 411 | 423 | 407 | 406 | 410 |
| 0.73 | 232 | 432 | 427 | 441 | 409 | 417 | 428 | 424 | 415 | 429 |
| 0.69 | 231 | 433 | 429 | 436 | 405 | -   | 428 | 443 | 420 | 418 |
| 0.66 | 235 | 439 | -   | 424 | 402 | -   | 433 | 415 | 422 | 417 |
| 0.63 | -   | -   | -   | 431 | -   | -   | 436 | 434 | 430 | 419 |
| 0.59 | -   | -   | -   | 427 | -   | -   | -   | 449 | 425 | 422 |
| 0.56 | -   | -   | -   | 441 | -   | -   | -   | 426 | -   | 424 |
| 0.54 | -   | -   | -   | 440 | -   | -   | -   | 430 | -   | -   |
| 0.51 | -   | -   | -   | 441 | -   | -   | -   | 431 | -   | -   |

We note that due to the very directional interactions of the patches in the 0.1 simulations, two clusters had formed. Throughout the simulation these clusters were constantly combining and breaking apart from each other, leading to the variation in the cluster size.
Table S5: Cluster sizes for phase determination in the window-to-arene simulations.

| $k_B T$ | 0.1 | 0.2 | 0.3 | 0.4 | 0.5 | 0.6 | 0.7 | 0.8 | 0.9 | 1.0 |
|--------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| 1.04   | -   | -   | -   | -   | -   | -   | 103 | 81  | 61  | 109 |
| 0.99   | -   | 434 | 436 | 424 | 437 | 182 | 245 | 226 | 188 | 175 |
| 0.94   | -   | 449 | 444 | 432 | 448 | 291 | 332 | 314 | 311 | 319 |
| 0.90   | -   | 448 | 439 | 430 | 440 | 337 | 356 | 351 | 338 | 283 |
| 0.85   | -   | 443 | 439 | 423 | 448 | 427 | 404 | 380 | 372 | 380 |
| 0.81   | -   | 445 | 442 | -   | -   | 436 | 421 | 401 | 378 | 386 |
| 0.77   | -   | 444 | -   | -   | -   | 440 | 422 | 420 | 417 | 422 |
| 0.73   | -   | -   | -   | -   | -   | 441 | 438 | 419 | 409 | 417 |
| 0.69   | -   | -   | -   | -   | -   | 433 | 426 | 426 | 425 |     |
| 0.66   | -   | -   | -   | -   | -   | -   | 446 | 427 | 430 |     |
| 0.63   | -   | -   | -   | -   | -   | -   | -   | 436 |     |     |
| 0.59   | -   | -   | -   | -   | -   | -   | -   | -   |     |     |
| 0.56   | -   | -   | -   | -   | -   | -   | -   | -   |     |     |
| 0.54   | -   | -   | -   | -   | -   | -   | -   | -   |     |     |
| 0.51   | -   | -   | -   | -   | -   | -   | -   | -   |     |     |

3 Additional structural images
4 Structural determination

Clusters formed from HOOMD-Blue simulations were transformed onto clusters of cages using custom made code. This code transformed for the distance between neighbouring octahedra from $\approx 0.6 \text{Å}$ as in the simulations, to $\approx 20 \text{Å}$ between the cages for easy visual examination of the structure. A second custom made code then picked out a unit cell of repeating cages (Fig. S2(a)). As the unit cell contained orientations of the cages which were slightly disordered due to the nature of the simulations, the structure was coarse-grained in order to use traditional symmetry solving algorithms. The coarse graining process worked by creating a structure with only the central position of the cage and the position of three of the carbons of the arene, in order to preserve the symmetry of the cage (Fig. S2(a-c)). Using this coarse-grained structure, $FINDSYM^{S1, S2}$ was employed to determine the structures space group. The same transformation as invoked by $FINDSYM$ to the coarse-grained unit cell, was then applied to the original unit cell (Fig. S2(e)). Using the solved space group, the symmetry conditions were then applied to the unit cell, and the sites that
related to the same atomic position were merged to get the full structure of the cage under symmetry conditions. This process worked best for cages with a small patch width, but for some of the ordered phases at large patch widths for the window-to-arene simulations, the orientation of the coarse-grained structures were still too disordered to solve using this process, *i.e.* FINDSYMM was unable to find an ordered structure even with the coarse-grained configuration. In these cases, the clusters were instead compared visually to known solid-state structures of octahedral porous organic cages. An overview of which of the space groups of the window-to-arene structures were solved using the coarse-graining process and which were solved visually is given in Table S6.
Figure S2: Overview of the process used for finding the space group of the structures shown for the ordered window-to-window simulation when $\sigma_{\text{ang}} = 0.3$. (a) An example unit cell (shown in black) picked out from the simulated cluster with the positions of the cages neatened to be on likely high symmetry sites i.e. $\frac{1}{4}, \frac{2}{4}, \frac{3}{4}$ or $\frac{1}{3}, \frac{2}{3}, \frac{3}{3}$ etc. (b) Addition of a central atom to each cage (purple) and selection of three of the carbon atoms (yellow) on the arene chosen to preserve the symmetry of the cage whilst reducing the degrees of freedom. (c) The structure of just the central position and three arene carbon atoms used in FINDSYM to get the space group of the structure. (d) The outputted structure from FINDSYM with the space group F4132. (e) Structure obtained by converting the original unit cell to the same unit cell as outputted from FINDSYM with no symmetry operations. (f) Final structure obtained by applying the symmetry conditions of the space group onto the structure shown in (e) and merging the atoms that sit around the same crystallographic sites.
Table S6: Whether the structure was solved by eye or by findsym

| $\sigma_{\text{ang}}$ | Solved using coarse-graining and findsym |
|------------------------|------------------------------------------|
| 0.2-0.4                | Yes                                      |
| 0.5                    | Yes                                      |
| 0.6                    | No                                       |
| 0.7                    | No                                       |
| 0.8-1.0                | NA                                       |

Figure S3: Comparison of the simulated phases of the window-to-arene simulations (top) to their corresponding experimental phases (bottom) as solved visually. (a) $\sigma_{\text{ang}} = 0.6/\text{CC9}$, the experimental structure is derived from CC9 with the vertex functional groups (phenyl groups) removed for easy comparison. (b) $\sigma_{\text{ang}} = 0.7/\text{CC1}\beta$. 
5 Phase diagram determination

The phase diagrams for both the window-to-window and window-to-arene simulations were coloured using a continuous colour map based on their similarities to the RDF of the solved ordered structures at low temperature Fig. S6. For phases at small patch widths, the structures are relatively ordered and as such the RDFs were compared to the RDFs of the ordered structure calculated with 40 bins, *i.e.* a high resolution RDF (Fig. S6(a),(c)). On the other hand, the phases at large $\sigma_{ang}$ are inherently more disordered, and as such so are their RDFs Fig. S4,S5. Therefore, the similarities between the simulated and solved structure was compared using an RDF with a fewer number of bins (25) *i.e.* an RDF lower resolution.

The comparisons between the RDFs were made by using a time series analysis, dynamic time warping, an algorithm commonly used for measuring the similarity between two temporal sequences. This analysis has been used to compare the difference between two PXRDs before.$^5$ In this study, we used it to compare RDFs by first normalising the RDFs such that the largest value of $g(r) = 1$, and then calculating the dynamic time warping. The values returned by the algorithm were then normalised for both the window-to-window and window-to-arene simulation, which gave a similarity measure between 0-1 for each ordered RDF where 1 is the most similar and 0 the least. This similarity measure was then used to colour the phase diagrams. For the window-to-window simulation, given there was only one phase deemed to have both orientational and translational order, the phase diagram was coloured orange by the similarity measure to said phase (Fig. S6(a)). For the window-to-arene simulation, as there were four separate ordered phases, the similarities of the RDFs for the most disparate three were used to colour the phase diagram red, green, and blue (Fig. S6(b-d)).
Figure S4: The RDFs for the low temperature structures from the window-to-window simulations when (a) $\sigma_{\text{ang}} = 0.1$, (b) 0.2, (c) 0.3, (c) 0.4, (d) 0.5, (e) 0.6, (f) 0.7, (g) 0.8, (h) 0.9, and (i) 1.0.
Figure S5: The RDFs for the low temperature structures from the window-to-window simulations when (a) $\sigma_{\text{ang}} = 0.2$, (b) 0.3, (c) 0.4, (c) 0.5, (d) 0.6, (e) 0.7, (f) 0.8, (g) 0.9, and (h) 1.0.
Figure S6: The RDFs used for comparison when colouring the phase diagrams from the window-to-window and window-to-arene simulations. (a) The RDF used in the window-to-window simulation to colour the phase diagram orange (b) the RDF of the red component used to colour the window-to-arene phase diagram which corresponds to the phase where \( \sigma_{ang} = 0.7 - 1.0 \) (c) the RDF of the blue component used to colour the window-to-arene phase diagram which corresponds to the phase where \( \sigma_{ang} = 0.5 \) (d) the RDF of the green component used to colour the window-to-arene phase diagram which corresponds to the phase where \( \sigma_{ang} = 0.2 - 0.4 \).
6 Arene-to-arene configuration

Figure S7: Representative configuration for the arene-to-arene simulations where $\sigma_{ang} = 0.3$. 
7 High $\sigma_{\text{ang}}$ leads to disorder

Figure S8: (a) Cross section of a representative configuration for the window-to-arene simulations where $\sigma_{\text{ang}} = 0.9$ showing that the orientational behaviour becomes disordered at high $\sigma_{\text{ang}}$ and $\sigma_{\text{tor}}$. (b) Representative cluster of the amorphous behaviour at high temperature for simulations where $\sigma_{\text{ang}} \geq 0.8$. This configuration is taken from the simulation where $\sigma_{\text{ang}} = 0.7$. CIF files for both these phases are provided in the ESI.
8 Porous window-to-arene configurations

Figure S9: Representative configurations and 1D pore channels (shown in yellow) of the structures from the window-to-arene simulations where (a) $\sigma_{\text{ang}} = 0.2 - 0.4$, and (b) $\sigma_{\text{ang}} = 0.5$. 

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9  Predictive capabilities

To determine whether atomistic calculations can be used to determine the likely values of $\sigma_{\text{ang}}$ for known cages, we looked at how the energies of two cages slipping over one another changes as a function of position and distance. For this we focused on the energetics of CC9 and CC1 due to them both having solid-state structures which exist within the same simulations; window-to-arene. We first fully optimised both CC9 and CC1 in the mixed Gaussian and plane waves code CP2K/QUICKSTEP with the PBE functional, GTH-type pseudopotential, TZVP-MOLOPT basis sets, using a cutoff of 400 Ry for the plane wave grid and the Grimme-D3 dispersion correction. With the optimised cages, we than ran single point calculations using the OPLS3 force field as implemented using Schrödinger on two cages with the same orientation which were displaced next to each other along the (111) axis (Fig. S10(a)). To determine the preferred orientation and displacement between the two molecules, we rotated one of the cages around the (111) octant at different distances from the first cage (Fig. S10(b)). To monitor how the energy of the dimers changed with position, we translated one of the cages around the (111) octant at different distances from the first cage (Fig. S10(c)). For these calculations we constrained the vertices of the cages so as to ensure we were sampling the energy of the rigid cages rotating/slipping, rather than possible conformational effects. We then measured the interaction energy between the cages in each pair by calculating:

$$E_{\text{int}} = E_{\text{pair}} - 2E_{\text{cage}}$$

(1)

For the rotation calculations, we plotted all points on a graph where $E_{\text{int}} < 0$ (Fig. S11). These showed that the bulkier side groups considered here would not effect $\sigma_{\text{LJ}}$ and as such our octahedral shapes can be considered a crude representative of all the cages. For the slipping calculations, we then plotted all the energies where $(E_{\text{int}} - E_{\text{int}}^{\text{min}}) \leq 30 \text{ kJmol}^{-1}$ where $E_{\text{int}}^{\text{min}}$ is the minimum interaction energy between two cages (Fig. S12). We only looked...
at energies within 30 kJmol\(^{-1}\) as we deemed that energies higher than 30 kJmol\(^{-1}\) would be energetically unfavourable. These results showed that CC1 has more freedom in the slipping motion than CC9 as there are positions off the central \(\langle 111 \rangle\) axis which are within the 30 kJmol\(^{-1}\) limit for CC1, whereas CC9 only has on axis low energy configurations. This helps explain why CC9 is found at a lower value of \(\sigma_{\text{ang}}\) than CC1.
Figure S10: (a) Example of a configuration used in a dimer calculation with CC1 cages where the cages are displaced along the $\langle 111 \rangle$ axis, shown as a blue arrow. (b) Vector (shown in red) that the cage was rotated around and displaced along to examine the effect of rotation and displacement on the interactions between the two cages. (c) Octant (shown in blue) that the cage was displaced around to monitor the effect of cage slipping on the interaction energy.
Figure S11: Polar plots showing the energetics of cages rotating around and displacing along the ⟨111⟩ axis for (a) CC1 and (b) CC9. Here r is the distance along the ⟨111⟩ axis and θ is the angle of rotation. The colour of the plot is a measure of the interaction energy where the lower the energy, the darker it is.
Figure S12: The energetics of cages slipping over each other for (a) CC1 and (b) CC9. The left hand side is looking down the $<111>$ axis and the right hand side is a cross section containing the lowest energy position looking down the $z$ axis. Points are shown when the energy of the cages slipping at each position is $\leq 30$ kJmol$^{-1}$. The colour of the plot is how close to the minimum the energy is, where the closer to the minimum energy it is, the darker green it is, and the closer to the limit it is, the more yellow it is.
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