Research Roundup

A chaperone feels the heat

A chaperone is supposed to keep its cool when temperatures get hot. But Syed Rizvi, Laura Mancino, Malini Raghavan (University of Michigan, Ann Arbor, MI), and colleagues show that calreticulin—a glycoprotein chaperone—starts to melt in the heat. The resulting structural changes actually improve its chaperone activity and may also occur transiently under normal conditions.

Calreticulin is a stress-induced chaperone that helps glycoproteins such as MHC Class I molecules fold properly in the ER by binding to the target proteins' sugar groups. The new results show that at high temperatures or low calcium levels, calreticulin can also bind independently of sugars and thus more effectively inhibit protein aggregation.

Protein binding was accompanied by structural changes in calreticulin, including oligomerization at high temperatures. "It has remarkable conformational lability for a chaperone," says Raghavan. "Its stability is like [that of] the substrates themselves." The COOHterminal tail was found to be necessary to inhibit the oligomerized form, but its effect may be overcome at high temperature or low calcium so that the protein can help out when the ER is overwhelmed. JCB

Reference: Rizvi, S.M., et al. 2004. *Mol. Cell.* 15: 913–923.



Heat-shocked calreticulin (HS-CRT) is better able to prevent protein aggregation (purple bars).

Cells have a bubbly look

century ago, scientists noted a similarity between the patterns that cells and soap bubbles adopt. Now, this nearly forgotten parallel is recalled by Takashi Hayashi and Richard Carthew (Northwestern University, Evanston, IL). Their results show that the same physical mechanism that drives bubbles to coalesce in forms that minimize their total surface area (and thus their surface free energy) also governs cell patterning.

Like bubbles, cone cells in the fly eye assembled in shapes that minimize surface area. Parameters that help specify this minimization include the number of intersection points between cells or bubbles, the number of interfaces that link these points, and the angles between them. Ommatidia containing abnormal numbers of cone cells still retained these minimizing characteristics. But those that lacked cadherins, and thus intercellular adhesion, did not. Presumably the lack of adhesion makes these cells act as independently energy-minimizing units rather than a single unit.

As well as controlling cone cell packing, cadherins and surface mechanics also organize cone cells within an ommatidia. Only cone cells expressed N-cadherin; other ommatidial cell types, such as pigment cells, expressed E-cadherin. Cone cells thus aggregated in the midst of the other cells to mini-

mize contact between cells of different adhesion strengths (N-cadherin adheres more strongly than does E-cadherin). By altering the adhesion differential through misexpressed or mutant cadherins, the authors perturbed the shape of the cone cell group and thus increased its area of contact with other cells. JCB



Reference: Hayashi, T., and R.W. Carthew. **N** 2004. *Nature*. 431:647–652. **N**

Mutant cone cells (purple) that lack N-cadherin show patterning defects.

Uncooperative immune cells

t's hardly *The Art of War*. In what seems like a counterproductive effort, the immune system's first line of defense sets up a barricade to block the next wave of defenders, as shown by Ravi Rao, Francis Luscinskas (Harvard Medical School, Boston, MA), and colleagues.

The first immune cells to arrive at injured tissues are usually neutrophils. These cells transmigrate through the endothelial cells lining the vessel wall and release granules full of proteases that chew up the damaged tissue or infectious agents. Now, one of these proteases, called elastase, is shown to act on endothelial cells to deter the entry of later-arriving T cells.

Elastase hindered T cell transmigration by cleaving and inactivating the vessel-bound chemokine SDF-1 α , which guides T cell migration on the vessel wall to entry sites. "When we started," says Luscinskas, "we thought the neutrophils might enhance chemokine activities at the blood vessel wall, where more white blood cells are destined to transmigrate. But the opposite is true." By closing the door on T cells, neutrophils might prevent potentially dangerous overactive immune responses.

Although protease inhibitors are abundant in the blood stream, plasma did not improve T cell entry. Elastase may be protected from the inhibitors by the neutrophil cell body, which physically seals off plasma inhibitors from accessing the space between the neutrophil and endothelium. JCB Reference: Rao, R., et al. 2004. *J. Exp. Med.* 200:713–724.