RESEARCH NEWS

with it could be an important regulator of protein synthesis.

One way to interfere with the process would be to bind eIF-4E so that it is no longer free to join up with the mRNA. Sonenberg's group was looking for such 4E-binding proteins, and they had already found two, for which they cloned the genes. When they realized that one of these 4E-binding proteins was PHAS-I, "it was a pleasant surprise," says Sonenberg. "We knew it represses translation, but we couldn't link it to signal transduction."

Along with Lawrence's biochemical experiments, the link to eIF-4E activity established that insulin-activated MAP kinase phosphorylates PHAS-I at a particular site, which causes it to dissociate from eIF-4E, so that the factor can initiate translation. The PHAS-I work "sure is interesting," says Simon Morley of the University of Sussex, U.K., who also studies the regulation of initiation.

The idea that signals from growth factors

and hormones can be relayed via MAP kinase directly to the protein translation machinery, rather than traveling via the events in the nucleus, is just beginning to gain acceptance with researchers. But for some, the new results vindicate a long-held belief. Cohen comments, "I'm not surprised-we've been pushing the idea for a long time that MAP kinase has actions [outside the cell's nucleus]." Biochemist George Thomas of the Friedrich Miescher Institute in Basel, Switzerland, was one of the first scientists to realize the potential importance of such signaling pathways 20 years ago. He says he has "argued from the beginning" that mitogens would be involved in the regulation of translation. "It's great to have additional evidence of direct effects of mitogens on ... the translational machinery," he adds.

Although the two papers are based on the single case of fat cells stimulated by insulin, the presence of PHAS-I in other cell types

strongly suggests that it will play a key role in other cells and possibly in other pathways. Lawrence points out, "We don't know that MAP kinase is the only regulator [of PHAS-I]-such an important inhibitor is probably the target of other second messenger systems." Sonenberg is therefore busy examining the other phosphorylation sites on PHAS-I. And it's likely that PHAS-I, and the related protein coded by Sonenberg's second cDNA clone, are two members of a family of similar proteins. As Denton says, "This is an incredibly exciting time in protein translation." And, because of the link made in the Lawrence paper, that excitement is not limited to those interested in insulin-it's also spilling over into the fastgrowing field of growth factors.

-Claire O'Brien

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___NANOENGINEERING_

AFM Fabricates a Tiny Transistor

DAVOS, SWITZERLAND—Nanoengineering—the creation of structures smaller than 100 nanometers, or 0.1 micrometer—holds particular allure for the semiconductor industry, where smaller is always better. Until now, however, the promise has remained largely theoretical, because conventional chip-making techniques have trouble with anything smaller than 0.2 micrometer, and newer methods have been confined to the laboratory. But at a meeting here last month, Stanford University applied physicist Calvin Quate reported creating working transistors using a tool created for nanoscale observation, the atomic force microscope (AFM).

Many delegates believed this is the first example of a working device made with an AFM and, while just on the border of the nanoscale range, the fact that AFMs have the potential to make devices much smaller is getting people excited. "This is really new," says Jürgen Mohr of the Institute of Microstructure Technology at the Karlsruhe Nuclear Research Center in Germany. "We now see that it's possible to use the AFM for fabrication, not just observation."

Quate's report was a last-minute addition to the opening session of the Micro and Nano-Engineering '94 conference. Traditionally geared towards conventional chip-making techniques, the meeting broadened its scope for the first time this year to include local probe techniques such as the AFM. The result? A 70% hike in attendance and a jump in submitted papers from 100 to 172—nanofever was in the air.

The AFM and its sister instrument, the scanning tunneling microscope (STM), were originally designed for imaging surfaces with

atomic-level detail. Both microscopes rely on an extremely fine tip; in the AFM the tip, mounted on the end of a cantilever, rests on a surface. As the tip is moved horizontally, it traces the irregularities of the surface, and a sensor measures the up-and-down movements of the cantilever to generate an image. But in the past several years researchers have also been using them to modify surfaces: punching holes, carving grooves, depositing material, and causing chemical reactions.

Quate and his team used an AFM to draw an ultrafine line that forms the heart of a transistor. They started with a sapphire substrate and coated it with a layer of amorphous silicon and an overlayer of hydrogen. They then moved an AFM tip over the surface, sending a mild electric current from tip to surface. The current cleared away the protective hydrogen layer beneath the tip while oxidizing the surface of the silicon by activating water molecules in the surrounding atmosphere. The resulting silicon dioxide line protected the silicon beneath it while the rest of the silicon was chemically removed from the substrate. The team succeeded in drawing lines less than 0.1 micrometer wide.

Using this AFM line-drawing technique, the team made a number of devices called metal-oxide-silicon field effect transistors (MOSFETs). In such devices, a current passing through a piece of semiconductor is controlled by an electrode above it called the gate. The shorter the gate the faster the transistor. The Stanford team made the bulk of each transistor using conventional lithography, where successive layers are built up and then parts etched away using a series of masks. The AFM was used only for making



Starting line. A probe microscope was used to make an electrode $0.1 \mu m$ wide on this transistor.

the gate. (The line width corresponds to the gate length.) Twelve of the transistors they made, with gate lengths ranging from 0.7 to 0.2 micrometer, behaved like conventionally made MOSFETs in performance tests.

Older techniques can create transistor gates just as small as those made by Quate's team, but while conventional lithography is pushing up against the physical limits of miniaturization, scanning probe techniques are still in their infancy and have the potential to go much smaller. This prospect is now drawing interest from industry, but the process is at the moment far too slow for production: Local probes can draw a line on only one device at a time, while conventional lithography imprints millions of transistors in one process. To speed things up, Quate envisions an array of up to 10,000 AFM tips working simultaneously. His team has been experimenting with a prototype five-tip array. Another advantage is cost: Local probes are cheaper than conventional equipment.

-Elizabeth Gardner

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