

The eight biospherans were selected and trained, not as scientists, but as resourceful individuals with survival skills who could work together to live for 2 years at a subsistence level. In reality, Biosphere 2 is research at the interface between the natural and the social sciences, where the real world problems of the future lie.

Overall, the mission of Biosphere 2 is to find out if approximately 1 hectare of pollution-free landscape (that is, with no industries or automobiles) consisting of 80% natural or seminatural ecosystems (the biomes) and 20% labor-intensive polyculture agriculture would provide bioregenerative life support for eight people. The answer is apparently going to be, "just barely." The experiment could also be considered a microcosm test of the Gaia Hypothesis—that an adequate flow of high-quality energy (sun, electricity, and natural gas, in this case) and a diversity of life forms will co-evolve or self-organize into a system that will support life. It is inconceivable to me that anyone would expect an instant achievement of balances, such as between oxygen and carbon dioxide, when we don't fully understand how such balances are maintained in Biosphere 1 (the Earth).

When one considers that nothing on the scale of Biosphere 2 has been attempted before (NASA's designs for regenerative life support are entirely different, and much smaller) and how little we really know about how our Biosphere 1 works, a measure of success will have been achieved if the biospherans come out alive and healthy this fall after the 2-year isolation. Certainly the experiment will have improved our understanding of human-biosphere interrelations and helped answer the question of how much natural environment must be preserved for life support, and it will have provided a basis for improving the design next time around.

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Judging Science

In his informative article on the *Daubert et al. v. Merrell Dow* case now before the U.S. Supreme Court ("Supreme Court to weigh science," *News & Comment*, 29 Jan., p. 588), Eliot Marshall notes that the American College of Legal Medicine (ACLM) has filed a friend-of-the-court brief on behalf of Merrell Dow, asking that the lower court decisions be upheld. The Carnegie Commission brief Marshall discusses is not the only one that "proposes a method for screening

scientific testimony"; ACLM's brief does so as well. We ask the court to consider the following questions: Was a controlled study performed? Were the results statistically significant? Has the study been published in a peer-reviewed journal? The evidence proffered in this case did not meet all of those criteria. Thus, we believe it did not attain the threshold that would permit it to be helpful to a jury, and the courts were correct to exclude it under the federal rules.

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Marshall's article about the upcoming *Daubert* case is valuable because it goes beyond the details of a specific case and presents the broader issues of scientific standards in the courtroom. I want to clarify two points raised in this article that arise because of my involvement in this case.

After describing the plaintiffs' analyses, Marshall writes, "And, so far, it hasn't been presented to a jury, because the case has been dismissed in each courtroom . . ." On the contrary, this evidence was admitted by the court in at least five cases involving the antinauseant drug Bendectin.

Marshall correctly states that I direct "a California state health department group that monitors reproductive risk." However, the arm of the department that is devoted to the study of birth defects is distinct from this group. Furthermore, the analysis of Bendectin that I performed was not carried out as part of my work for the state, and does not represent the views of the department but are my own.

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Multidrug Resistance-Associated Protein: Sequence Correction

In the report "Overexpression of a transporter gene in a multidrug-resistant human lung cancer cell line" (4 Dec. 1992, p. 1650) (1), we presented the predicted amino acid sequence of a novel adenosine triphosphate-binding cassette transporter gene that we designated "multidrug resistance-associated protein" or "MRP."

We recently discovered a typographical error in the nucleotide sequence of MRP that resulted from the introduction of an

		10	20
Original		MAPTRSGTGMSRGIPATPSP	
Corrected	MALRGFCSDGSDPLWDNVNTWNTSNPDFT		
	10	20	30
Original	30	40	50
	SAFRTSSCGCLVFTSGPVPFFYFLYLSRH		
Corrected	KCFQNTVLVWVPCFYLWACFFPYFLYLSRH		
	40	50	60

Fig. 1. Corrected MRP-deduced amino acid sequence.

additional thymidine residue at position 206 of the original DNA sequence. This additional thymidine residue is not present in any of our original sequencing autoradiographs, partial sequence files, or the original compilation of the sequence. It was introduced as result of a typographical error during the manipulation of the sequence to generate forms suitable for publication and appears in our report.

The additional base in the nucleotide sequence renders incorrect the predicted sequence of the first 40 amino acids of the translated open reading frame. As a consequence, the predicted size of MRP is 1531 amino acids rather than 1522, as we stated. The correct sequence appears in Fig. 1 and has been corrected in GenBank.

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References

1. S. P. C. Cole *et al.*, *Science* 258, 1650 (1992).

Knowing It All

An "omniscient super-physicist" who did not care to measure "the velocity, momentum, and every other property of every particle in the universe" (*Research News*, 26 Mar., p. 1824) could also "find out who killed JFK and how the dinosaurs died. . . ." In fact, being omniscient, she or he would not have to find out, but would know already.

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Corrections and Clarifications

The x-axis of the graph accompanying Jon Cohen's *News* article "Keystone's blunt message: 'It's the virus, stupid'" (16 Apr., p. 292) should have been labeled "Monkeys."

The footnote on page 482 accompanying Jon Cohen's *News & Comment* article "Drug companies join forces in search for AIDS therapy" (23 Apr.) should have listed Bristol-Myers Squibb Company as one of the participating companies in the collaboration.