data from statistically stationary processes where the record is much longer than the longest wavelength of interest. Most fossil time series violate all three constraints. The fossil record is our only possible data source for studying the long-term dynamics of Earth’s biotic system, but such studies require carefully designed hypothesis tests.

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Arsenic poisoning of Bangladesh groundwater

In Bangladesh and West Bengal, alluvial Ganges aquifers used for public water supply are polluted with naturally occurring arsenic, which adversely affects the health of millions of people. Here we show that the arsenic derives from the reductive dissolution of arsenic-rich iron oxyhydroxides, which in turn are derived from weathering of base-metal sulphides. This finding means it should now be possible, by sedimentological study of the Ganges alluvial sediments, to guide the placement of new water wells so they will be free of arsenic.

As many as a million water wells drilled into Ganges alluvial deposits in Bangladesh and West Bengal may be contaminated with arsenic1–6. Measured arsenic concentrations1–6 reach up to 1,000 μg l−1, which is above the limit set for drinking water in Bangladesh (50 μg l−1) or that recommended by the World Health Organization (10 μg l−1). Consumption of this contaminated water has led to widespread death and disease3–6.

Arsenic has been reported to derive from the oxidation of arsenic-rich pyrite in the aquifer sediments as atmospheric oxygen invades the aquifer in response to a lowering of the water level by abstraction7–9. However, this explanation is not consistent with the following observations7, made on

46 wells, typical of those in Bangladesh, that were sampled during May and June of 1997: in oxic (shallow) wells, arsenic concentrations are mostly below 50 μg l−1; in anoxic waters, arsenic concentrations (≤260 μg l−1) correlate with concentrations of dissolved iron (≤29 mg l−1) and bicarbonate (Fig. 1a); and arsenic concentrations increase with depth in wells at Manikganj, Faridpur and Tungipara. These observations suggest that arsenic is released when arsenic-rich iron oxyhydroxides are reduced in anoxic groundwater8, a process that solubilizes iron and its absorbed load and increases bicarbonate concentration. Sedimentary iron oxyhydroxides are known to scavenge arsenic9 and, in Ganges aquifer sediments, concentrations of diagenetically available iron (≤3.7%) and arsenic (≤26 pp.m) correlate well (Fig. 1b).

The arsenic-rich groundwater is mostly restricted to the alluvial aquifers of the Ganges delta10. The source of arsenic-rich iron oxyhydroxides must therefore lie in the Ganges source region upstream of Bangladesh. Weathered base-metal deposits are known to occur11–13 in the Ganges basin (at Bihar, Uttar Pradesh, West Bengal), so weathering of these arsenic-rich base-metal sulphides must have supplied arsenic-rich iron oxyhydroxide to downstream Ganges sediments during Late Pleistocene–Recent times. The arsenic-rich iron oxyhydroxides are now being reduced, causing the present problem. Reduction is driven by concentrations of sedimentary organic matter of up to 6%.

A knowledge of the sedimentary architecture and distribution of iron, arsenic and redoxt carbon in Ganges alluvial sediments will allow the development of a predictive model to guide future aquifer development in a way that minimizes arsenic pollution. Furthermore, as dissolved iron is oxidized it precipitates as iron oxyhydroxide, which scavenges arsenic from solution. It follows that simple aeration of anoxic Bangladesh groundwater, followed by settling, should remove a considerable amount of arsenic from solution. This simple treatment could be performed on a household or village scale. Although the disposal of the arsenic-rich iron oxyhydroxides would require special arrangement, this would be preferable to either the widespread poisoning that now exists or a return to the use of contaminated surface water for public consumption.

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Figure 1 Arsenic chemistry of groundwater and aquifer sediment from Bangladesh. a, Arsenic correlates with dissolved iron and with bicarbonate concentrations above 300 mg l−1; the latter is a by-product of iron oxyhydroxide reduction. b, Relation of arsenic and iron in aquifer sediments; arsenic concentrations are up to 15 times higher than crustal abundance.