

Arsenic in groundwater in the Bengal Delta Plain: slow poisoning in Bangladesh

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Abstract: The purpose of this paper is to provide an overview of the problems concerning the widespread occurrences of arsenic in groundwater in Bangladesh, a land with enormous resources of precipitation, surface water, and groundwater. Because of the potential risk of microbiological contamination in the surface water, groundwater was relied on as an alternate source of drinking water. Exploitation of groundwater has increased dramatically in Bangladesh since the 1960s to provide safe water for drinking and to sustain wetland agriculture. The presence of arsenic in the groundwater at elevated concentrations has raised a serious threat to public health in the region. Nearly 60–75 million people inhabiting a large geographical area are at potential risk of arsenic exposure, and several thousands have already been affected by chronic arsenicosis. The source of arsenic in groundwater is geogenic and restricted within the Holocene sedimentary aquifers. Mobilization of arsenic from the alluvial aquifers is primarily effected through a mechanism of reductive dissolution of the iron oxyhydroxides within the sediments, rather than by the oxidation of pyrite, as has been hypothesized by other workers. The problem is further accentuated by the fact that arsenic is also found at elevated concentrations in vegetables and rice grown in the areas where high-arsenic groundwater is used for irrigation. Dietary habits among the population are also an important pathway for arsenic ingestion. Studies are in progress at national as well as international levels to alleviate the arsenic crisis in Bangladesh. Besides the identification of arsenic-free tubewells in the affected areas for drinking purposes, purification of groundwater at household level by low-cost arsenic removal techniques is suggested. Rehabilitation of the patients with chronic arsenicosis and arsenic education programs for rural communities must be addressed urgently by the government of Bangladesh.

Key words: arsenic, groundwater, chemistry, redox, causes, effects, Bangladesh.

Résumé: L'objet de cet article est de proposer un panorama des problèmes résultant de la concentration d'arsenic dans les eaux souterraines au Bangladesh, une terre riche en eaux de précipitation, de surface et souterraines. Vu le risque potentiel de contamination micro-biologique de l'eau de surface, l'eau de nappes profondes s'est révélée être une alternative pour offrir de l'eau potable. L'exploitation des nappes a augmenté au Bangladesh de manière dramatique à partir des années 60, afin de fournir une source sûre d'eau potable et de soutenir l'arrosage agricole. La présence d'arsenic à concentration élevée dans les nappes augmente les menaces sérieuses pour la santé publique de la région. Environ 60 à 75 millions

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de personnes sont soumises à un risque potentiel de contamination à l'arsenic et plusieurs milliers de personnes ont déjà été affectées par de l'arsenicose chronique. L'arsenic dans les nappes est de source géogénique et se limite aux aquifères sédimentaires dans l'holocène. La mobilisation de l'arsenic des aquifères alluviaux est surtout dû à un mécanisme de dissolution réductrice du fer oxyhydroxides dans les sédiments, plutôt que d'une oxydation de la pyrite, hypothèse avancée par d'autres auteurs. Le problème est accentué par le fait que l'on trouve de l'arsenic à concentration élevée dans les légumes et dans le riz cultivés dans les zones irriguées avec de l'eau de nappe en teneur élevée d'arsenic. Les habitudes diététiques au sein de la population offre aussi une voie importante d'ingestion de l'arsenic. Des études sont en cours, tant au niveau national qu'international, afin de contrôler la crise de l'arsenic au Bangladesh. Par ailleurs, outre l'identification de puits potables dans la zone affectée, la purification de l'eau de sol dans les ménages par des techniques à faible coût est suggérée. La réhabilitation des patients atteints d'arsenicose chronique, et des programmes d'éducation/sensibilisation à l'arsenic pour les communautés rurales s'imposent d'urgence pour le gouvernement du Bangladesh.

Mots clés : arsenic, eau de nappe, souterraine, redox, causes, effets, Banglades.

[Traduit par la Rédaction]

1. Introduction

Bangladesh is situated in one of the world's largest deltas, located north of the Tropic of Cancer between 20°N and 26°N and 88°E and 92°E (Huq and Rahaman 1994). The country has common land borders with India to the north, west, and east and Myanmar to the south-east. To its south is the Bay of Bengal. The total surface area of Bangladesh is about 144 000 km² (55 126 square miles, comparable to the size of Wisconsin, U.S.A.) with a population of more than 120 million. It is one of the most densely populated countries in the world, with a population density estimated at 920 people km² of land and an annual population growth rate of 1.59% (1999).

The Bengal Delta Plain (BDP) (also known as the GBM Delta) is drained by three of the major rivers of the world — the Ganges, Brahmaputra, and Meghna (GBM) — which originate in the Himalayas and flow through India, Nepal, and China before flowing into Bangladesh and finally into the Bay of Bengal. Ninety-three percent of surface water in Bangladesh flows from adjoining countries, and precipitation (1.5–2.5 m) within the country accounts for only the remaining 7% of freshwater. More than 200 rivers and streams drain through Bangladesh, with a mean annual flow of water of about 38 000 m³ s⁻¹ (Reiman 1993; Huq and Rahaman 1994), carrying over 2.4×10^9 t of sediments each year before discharging finally into the Bay of Bengal (Khan 1994).

According to the statistics presented by Black (1990), there were about 50 000 tubewells in the erstwhile East Pakistan (currently Bangladesh) after the political division in the Indian subcontinent in 1947. The source of community water supplies in Dhaka and other regional cities in Bangladesh was groundwater, mainly abstracted from the Dupi Tila aquifer. In other suburban areas and in most rural Bangladesh, surface water (mainly from ponds, locally known as *pukur*) and water from tubewells and (or) dugwells (shallow wells) were used for drinking and other household purposes. At that time, sufficient groundwater was not abstracted either for drinking purposes or irrigation because of poor infrastructure. Although Bangladesh is known to be a land of rivers, wetlands, and ponds, there has always been a scarcity of reliable fresh surface water free of bacteriogenic contaminants.

Starting in the 1960s, several technological development programs were initiated in Bangladesh by international agencies, including UNICEF, the World Bank, and others. More than 4.5 million shallow and deep tubewells were drilled in the country, with an aim to provide safe drinking water to about 97% of the population. The technology was economic at a minimal cost of US\$0.30 per metre (Black 1990). However, no water samples from these wells were analyzed for basic chemical quality. The success of the UNICEF-sponsored tubewell program resulted in reduced infant mortality, and reports on diarrheal dehydration of children under five declined from 250 000 in 1983 to 110 000 in 1996 (Anon 2000a). But since 1993 this program has become a matter of anxiety because of the manifestations of chronic

arsenic toxicity among the population in the entire region, resulting from prolonged consumption of groundwater. Recent studies have indicated that chronic arsenic toxicity is caused by consumption of arsenic naturally occurring in groundwater (Chakraborty and Saha 1987; Guha Mazumder et al. 1988; Saha and Chakrabarti 1995; Saha 1995; Mandal et al. 1996; Dhar et al. 1997, Roy Chowdhury et al. 1999).

At the end of the 20th century, the arsenic contamination in groundwater in Bangladesh has been documented as a serious environmental health disaster with severe socioeconomic consequences; a great challenge for the Government of Bangladesh is to provide safe drinking water for the population.

The present paper provides an insight into the historical background to the problem of safe drinking water in Bangladesh from surface water, groundwater, and rainwater sources. It includes special reference to the occurrences and causes of arsenic contamination in groundwater in the BDP aquifers of Bangladesh, which has emerged as a major crisis of the present decade. In addition, social problems due to the presence of arsenic in the drinking water and the present status of technologies available for the removal of arsenic are briefly discussed.

2. Water resources in Bangladesh

Bangladesh is essentially a land of rivers, with nearly 54 rivers entering the country. The annual flow of surface water through Bangladesh is estimated to be ca. $1\,511\,000 \times 10^6 \text{ m}^3$, of which $360\,000 \times 10^6 \text{ m}^3$ originates from outside the country (Baurne 1999). The mighty rivers Ganges (Padma) and Bramaputra (Jamuna) contribute 40% and 50% of surface water resources. Thus, nearly 90% of surface water in Bangladesh flows from neighboring countries. Local precipitation (1.5–2.5 m) within the country accounts for the rest of the freshwater resources.

2.1. Surface water

Two million hectares (e.g., 20% of the surface area) of Bangladesh is occupied by rivers, canals, and ponds (*pukur*). Water from these surface water sources is known to be free from natural arsenic, but the water is generally highly polluted and not suitable for drinking purpose without treatment. Industries are few in Bangladesh, mostly situated near big cities and in suburban areas. The surface water bodies, fisheries, and farmlands are, by and large, affected by the untreated effluents from industrial facilities, sewage, atmospheric depositions of toxic particulates, and agrochemicals. Leaching of salts from soils as a result of irrigation is a further problem for surface and groundwater. Surface water treatment plants are, however, expensive and complicated to build and operate in rural areas (Bangladesh Agriculture Sector Review 1989).

2.2. Precipitation

The average rainfall in Bangladesh is about 2200 mm, of which 75% occurs from May to September. The rainwater is free from arsenic and microbiological contamination. Several studies have indicated that it is possible to collect and store rainwater for drinking and household purposes. Because of lack of adequate technology, however, rainwater harvesting is not normally practiced in Bangladesh except in some areas of the Chittagong hill tracts (Chowdhury et al. 1987; BGS/MML 1999a).

2.3. Groundwater

Nearly 80% of the population in Bangladesh lives in rural areas with low socioeconomic status and poor infrastructure. A major challenge has been the provision of safe drinking water in rural sectors. Because of population growth, increased cultivation of high-yield crop varieties to attain self sufficiency in food grain production, and industrial development, the government of Bangladesh recognized the need for groundwater development for irrigation and drinking water supply (Hamid 1993). During the 1980s, the government of Bangladesh initiated groundwater-based irrigation projects within the National

Water Plan, with aid from the World Bank and United Nations Development Programme (UNDP), to improve irrigation facilities and meet the rising demand for safe drinking water. The total available groundwater reserve is estimated to be about $63\,800 \times 10^6 \text{ m}^3$, equivalent to an irrigation potential of $3.9 \times 10^6 \text{ ha}$. At present, 60% of the irrigated area is served by groundwater, with the remaining 40% served by surface water and seasonal rainfall. Groundwater has been a reliable source of water for irrigation and for potable water for a large fraction of the population in rural Bangladesh for the last 30 years. The number of domestic hand pumps now exceeds that of irrigation wells in Bangladesh (Pitman 1993).

Because of increased withdrawal of groundwater in rural communities for drinking and irrigation purposes, especially for winter paddy cultivation, the problem of arsenic contamination has become acute in the BDP (Bhattacharya et al. 1997; Dhar et al. 1997). In recent years, the presence of arsenic at concentrations above the permissible drinking water standard (0.01 mg L^{-1} , WHO, and 0.05 mg L^{-1} , Bangladesh standard) has been reported in groundwater (Fig. 1) over an area of nearly $118\,012 \text{ km}^2$ in 52 districts of Bangladesh (Karim 2000).

3. Distribution of arsenic in nature

3.1. Abundance of arsenic in the crust of the earth and in sediments

Arsenic is a natural constituent of the crust of the earth, ranking 20th in abundance in relation to other elements. It is a metalloid and a member of the nitrogen family. It is tasteless and odorless. Concentrations of arsenic in major types of rocks range between 0.5 and $2.5 \mu\text{g g}^{-1}$ (NAS 1977; Taylor and McLenan 1985; Kabata-Pendais and Pendais 1992), and its concentration in soils depends on parent rock characteristics, anthropogenic activities, climate, organic and inorganic components, and redox potential of the soil (Yan-Chu 1994). Arsenic has a long history of human use because of its toxic and medical properties. In the orient, arsenical compounds were used in medicine 2000–3000 years ago (Ferguson and Gavis 1972).

Arsenic is concentrated in minerals, either as arsenides of copper, lead, silver, or gold or as sulfides and sulfates. Major arsenic-containing primary minerals are arsenopyrite (FeAsS), realgar (As_4S_4), and orpiment (As_2S_3). Arsenic usually occurs in oxidized forms in sediments. In sandstones, the arsenic content varies from 0.6 to 120 mg kg^{-1} , whereas in shales and clay formations the concentrations may be as high as 490 mg kg^{-1} (NAS 1977). This higher value of arsenic in shales and clays is due to its accumulation during weathering and translocation in colloid fractions (Yan-Chu 1994). Typical concentration ranges of arsenic in crustal rocks are presented in Table 1. Concentration of arsenic varies enormously among coals from different sources, and its concentration range worldwide is reported to be between 0.5 and 80 mg kg^{-1} (average 10 mg kg^{-1}) (Clarke and Sloss 1992), although higher values have been reported occasionally.

In aquatic environments, inorganic arsenic occurs as oxyanions, namely in pentavalent arsenate ($\text{H}_n\text{AsO}_4^{n-3}$) and trivalent arsenite ($\text{H}_n\text{AsO}_3^{n-3}$) forms. Arsenate usually predominates under oxidizing conditions (Robertson 1989), whereas arsenite is predominant under reducing conditions, especially in sulfidic and methanic water (Korte and Fernando 1991; Pokrovski et al. 1996). Arsenate is commonly adsorbed onto the surface of iron, aluminum, and manganese oxides under an oxidizing environment and hence is immobile. Under reducing conditions transformation of arsenate into arsenite facilitates mobilization of arsenic from the sediments to the groundwater. However, the solubilities of arsenic species play an important role in controlling its pathways in the environment (Thornton 1999).

3.2. Pathways of arsenic emission

Arsenic input into the ecosystem can be divided into two categories: natural and anthropogenic emissions. There are several reports that cover both categories of arsenic emissions to the atmosphere on global, regional, and local scales (Nraigu and Pacyna 1988; Nriagu 1989; Pacyna 1996, 1998). Once

Table 1. Natural abundance of arsenic in crustal materials (Jacks and Bhattacharya 1998).

Rock type	Abundance (mg kg ⁻¹)
Igneous rocks	
Ultrabasics	0.3–16
Basalts	0.06–113
Andesites	0.5–5.8
Granites/silicic volcanics	0.2–13.8
Sedimentary rocks	
Shales and clays	0.3–490
Phosphorites	0.4–188
Sandstones	0.6–120
Limestones	0.1–20
Coal	1.0–1500
Peat	16–340

it is introduced into the atmosphere, arsenic can circulate for a long period of time before transfer to a permanent sink. Soluble (reduced) forms of arsenic in water tend to be quite mobile, whereas less soluble forms tend to adsorb onto sediments and clays or other mineral soil components. Microbial activity in soils, sediments, and water produce the methylated forms of arsenic that are volatile in nature and can re-enter the atmosphere and ultimately change back into inorganic forms.²

3.2.1. Industrial emissions

Information on anthropogenic release of arsenic to the environment of Bangladesh is not available. There are about 30 000 industrial facilities in and around the major cities where soil arsenic concentrations mostly range between 2.13 and 4.27 mg kg⁻¹. However, in the vicinity of industrial sites some hot spots with soil arsenic concentrations as high as 5020 mg kg⁻¹ have been recorded (Ullah et al. 1999). As for the energy sector, total production was about 22.44 Mtoe (million tonnes of oil equivalent) from oil, natural gas, hydroelectricity, and combustible renewable wastes. In Bangladesh, there are no coal-fired thermal power plants. Diffuse emission of arsenic is expected to occur during the burning of wastes, jute, and oil and also from other high temperature processes.

In developed and developing regions industrial sources of arsenic generally include coal-fired power plants, smelting and melting of metals, incinerations of wastes, and the use of fertilizers, pesticides, chemicals, and solvents. Because of strict regulation, better abatement technologies, and process change, arsenic emissions to the atmosphere of Europe were reduced from 11 500 t per year (in the period 1960–1965) to 4 570 t in 1985 (Pacyna 1996; UN/ECE 1999). Future estimated projected arsenic emissions for Europe to be about 1900 t in 2000 (Pacyna 1998).

3.2.2. Natural sources

Natural emissions of heavy metals are affected by interaction of the atmosphere with the lithosphere, oceans, and cosmic space. In view of the numerous different pathways for the release of arsenic to the atmosphere, it is rather difficult to estimate the amount of this metalloid released, although Nriagu (1989) estimated global natural emissions of arsenic for 1983 to be 12 000 t (Ryaboshapko et al. 1999).

Weathering and leaching of arsenic-rich geological formations and mining wastes result in elevated concentrations of arsenic in natural waters in several parts of the world. The risk of arsenic contamination

² Ren, X. 1993. Study of environmental arsenic pollution from smelter and human exposure level in Yuman province. Paper presented at National Symposium on Environmental Criteria, Nanking, China, May.

is, however, much higher in groundwater than in surface water such as rivers, lakes, and reservoirs. Incidences of natural occurrence of arsenic in groundwater ($>10 \mu\text{g L}^{-1}$) are reported from several countries in the world, namely the United States (Welch et al. 2000), Canada (Grantham and Jones 1977), Argentina (Smedley et al. 1998; Bundschuh et al. 2000), Mexico (Cebrian et al. 1983; Armienta et al. 1995, 1997, 2000; Rodriguez et al. 2000), Chile (Borgoño and Greiber 1972; Borgoño et al. 1977), Ghana (Smedley 1996), Hungary (Varsanáyí et al. 1991), and Finland (Idman 1996). In Asia, arsenic-laden groundwater is reported in Taiwan (Lu 1990; Chen et al. 1994), China (Lianfang and Jianzhong 1994; Luo et al. 1997), Japan (Terade et al. 1960), southern Thailand (Choprapawon and Rodcline 1997), West Bengal, India (PHED 1991, 1993; Chatterjee et al. 1995; Bagla and Kaiser 1996; Bhattacharya et al. 1997), and most recently from Bangladesh (Dhar et al. 1997; Nickson et al. 1998; von Brömssen 1999). Table 2 summarizes the salient aspects of the distribution, concentration, and the possible mechanism for the release of arsenic in groundwater in a few affected countries.

4. Arsenic in the environment of Bangladesh

4.1. Review of the causes of arsenic contamination

Several factors have been assumed to have caused the substantial input of arsenic in the aquatic environment of Bangladesh (BGS/MML 1999a). These are (i) wooden poles treated with arsenic-based compounds supporting electric wires; (ii) uses of pesticides and chemical fertilizers; (iii) release of untreated effluent from facilities; and (iv) arsenic accumulated (enriched) in sediments by geological processes. Among the factors listed above, the first three are not directly responsible for the prevalence of widespread arsenic contamination in groundwater. However, these sources may lead to considerable contamination of soil and groundwater system elsewhere (Welch et al. 2000; Bhattacharya et al. 2001). Input of arsenic to agricultural soils may have been caused by the application of monosodium methanearsonate (MSMA) and lead arsenate (PbHAsO_4) as pesticides prior to the introduction of DDT (Welch et al. 2000). The most acceptable hypothesis, according to the present status of our knowledge, is that the cause of widespread arsenic contamination in the groundwater of BDP in Bangladesh is geological. Arsenic in the groundwater is released primarily from the sediments deposited during the Holocene period (Bhattacharya et al. 1998a, 1999a; BGS/MML 1999a).

4.2. Nature of the alluvial sediments in the Bengal Delta Plain

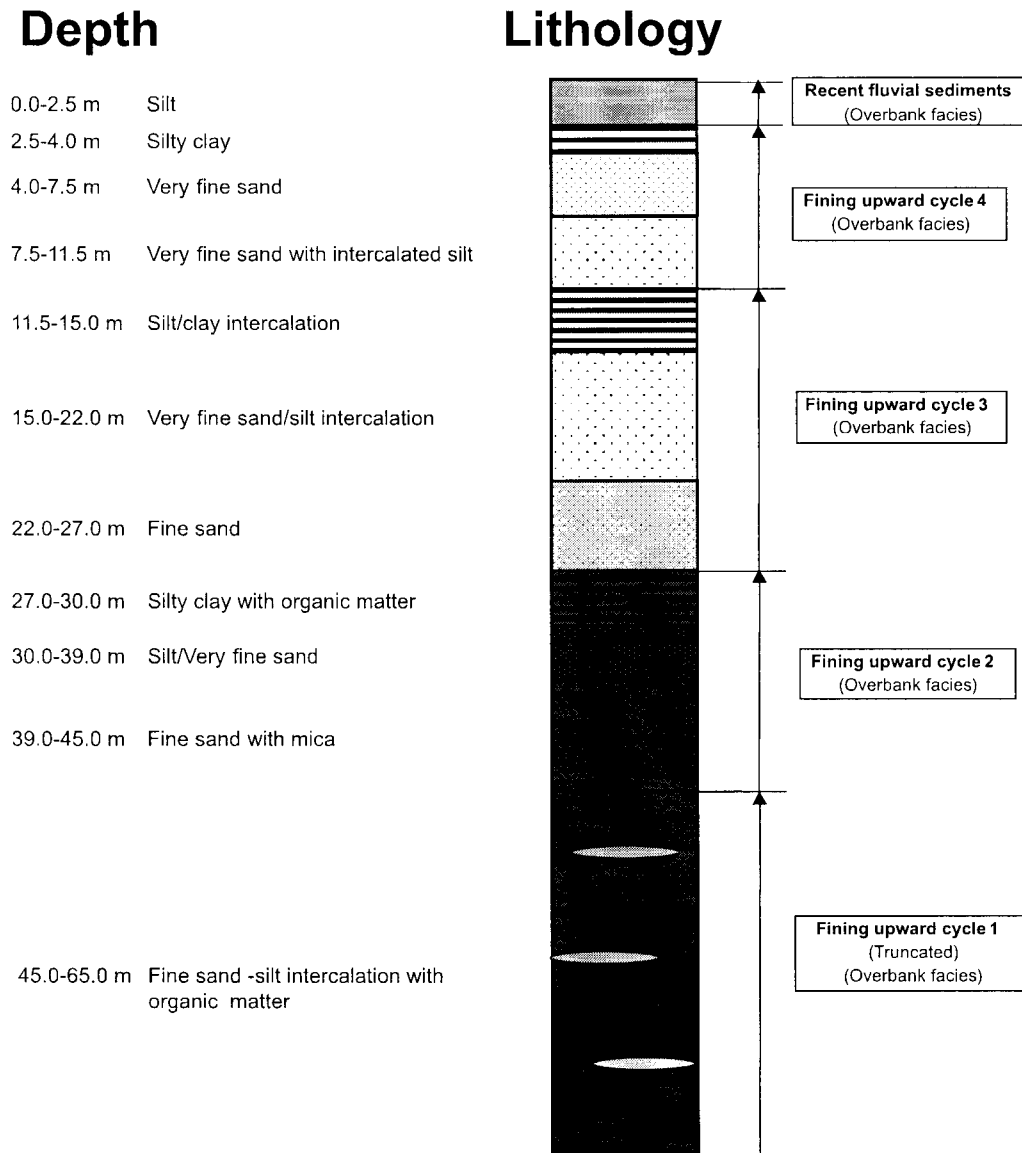
The sedimentary succession in the BDP consists of a thick sequence of sediments deposited by the Ganges, Brahmaputra, and Meghna rivers and their tributaries and distributaries (Bhattacharya et al. 1997, 1999a). The pool of arsenic-contaminated groundwater is hosted by the sediments deposited by the meandering river channels during the late Quaternary or the Holocene age. Lithology of these sedimentary successions include sand, representing the channel facies, and the overbank facies, comprising predominantly silt and clay, and exhibit a typical fining upward character (Bhattacharya et al. 1997, 1999a; Ahmed 1999). The fine-grained overbank facies are rich in organic matter (Nickson et al. 1998; Bhattacharya et al. 1999a; Mazumdar 2000). A typical sedimentary succession with stacks of fining upward sequences in Tala, Satkhira district in southwestern Bangladesh, is presented in Fig. 2.

The arsenic in these Holocene BDP sediments can be attributed to the parent rocks rich in arsenic from both the Himalayas and the highlands in the neighboring areas. Several isolated high arsenic geological domains are recognized that can be potential sources of the arseniferous sediments (BGS/MML 1999b; Bhattacharya et al. 1999a; Acharyya et al. 1999). These include (i) Samthan Hill near Darjeeling, West Bengal, India, sulphide minerals with arsenic concentration as high as 7.5%; (ii) arsenic-bearing nickel ores at Singbhum in Bihar, India; (iii) pyrite and chalcopyrite occurring widely in the granites and granitoid rocks of the Chottonagpur and Shillong Hills, India; (iv) iron ore deposits in eastern India are a major source of arsenic from the upper to the lower Ganges Bengal basin; (v) Gondwana coal

Table 2. Comparison of arsenic occurrences in groundwater from selected parts of the world.

Country/Region	Area affected (km ²)	Depth of well (m)	Arsenic conc. (g L ⁻¹)	Mechanism of contamination	References
Bangladesh, BDP (52 districts)	118 012	8-260	<2 to >900	Reduction of Fe-oxyhydroxides/Sulfide oxidation(?) in alluvial sediments?	Dhar et al. 1997; von Brömssen 1999; Karim 2000
West Bengal, India, BDP (8 districts)	34 000	14-132	<1 - 1300	Reduction of Fe-oxyhydroxides/Sulfide oxidation (?) in alluvial sediments	von Brömssen 1999; Bhattacharya et al. 1997; Mandal et al. 1996
China, Inner Mongolia (Huhhot Basin)	3 600	Shallow/deep	<100 - 1860	Reducing environment in alluvial sediments	Lianfang and Jianzhong 1994; Luo et al. 1997
Taiwan	---	Deep	Up to 1800	Oxidation of pyrite in mine tailings	Lu 1990; Chen et al. 1994
Thailand	10 districts	Shallow	120 - 6700	Oxidation of mine wastes and tailings	Choprapawon and Rodcline 1997
Ghana	1 600	70-100	2 - 175	Oxidation of arsenopyrite in mine tailings	Smedley 1996
Argentina (Chaco-Pampean Plains)	10 000 000	Shallow aquifers	100 - 4800	Volcanic ash with 90% rhyolitic glass	Smedley et al. 1998; Bundschuh et al. 2000
Chile	---	Shallow and deep wells	100 - 1000	Volcanic ash	Borgoño and Greiber 1972; Borgoño et al. 1977
Zimapan, Mexico	---	Shallow and deep wells	300 - 1100	Oxidation of sulfide from mine wastes	Armienta et al. 1997, 2000
Hungary (Great Hungarian Plain)	4 263	80-560	25 - >50	Complexation of arsenic with humic substances	Vársanayi et al. 1991
U.S.A.	518	53-56	100 - >500	Desorption of arsenic from Fe-oxyhydroxides/Sulfide oxidation	Welch 2000; Welch et al. 2000
Canada (Nova Scotia)	---	8-53	18 - 146	Oxidation of sulfides	Grantham and Jones 1977

Fig. 2. Succession of Holocene aquifer sediments with fining upward character of the deposits of meandering channels at Tala, Satkhira district in southwestern Bangladesh (redrawn from Mazumdar 2000).

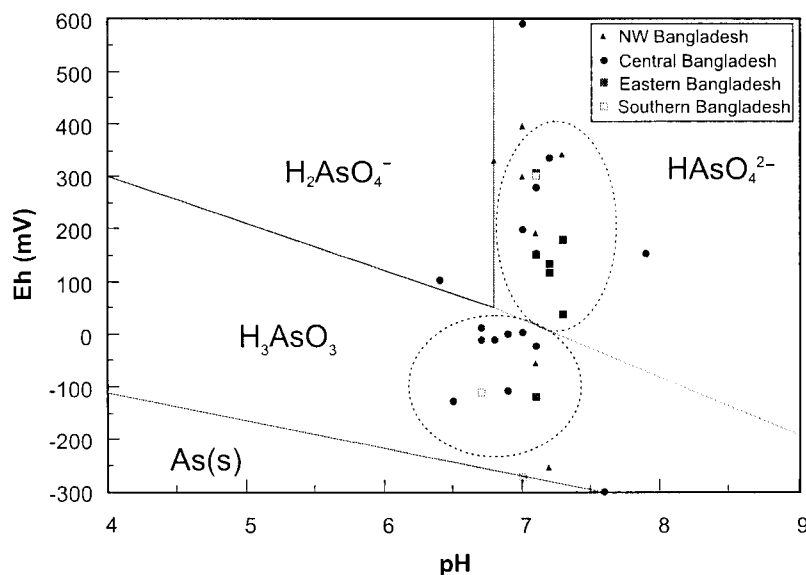


contains significant quantities of arsenic; (vi) volcanic terrain in Rajmahal (district Bihar), India; and (vii) the arc of the ultramafic suite of rocks occurring along the Indus Suture Zone. Another speculation about the potential source of arsenic is the Toba Ash, which resulted from the volcanic eruptions in the Indonesian archipelago during the Quarternary period.³

Thus the arsenic in the BDP sediments can be attributed to source terrains in the close proximity of the depositional basin. Arsenic liberated because of the oxidation of the primary arsenic-bearing sulfide minerals is brought to the BDP by the phases adsorbed onto suspended particulates, where arsenic is mostly adsorbed and co-precipitated with secondary Fe, Al, and Mn phases in the form of As(V)

³ Aftab Alam Khan, Bangladesh, personal communication.

Fig. 3. Eh–pH diagram showing the thermodynamic stability fields of trivalent As (H_3AsO_3 as unchanged species) and pentavalent As ($\text{H}_2\text{AsO}_4^{2-}$ and H_2AsO_4^- as the protonated species) along with the plot for the water samples from Bangladesh.



and finally deposited in the delta depending on the topography of the basin (Bagla and Kaiser 1996; Bhattacharya et al. 1997; Dhar et al. 1997; Nickson et al. 1998, 2000; Sahu et al. 2000).

4.3. Geochemical characteristics of arsenic in the Bengal Delta Plain groundwater environment

4.3.1. Groundwater chemistry and relation to arsenic

Groundwater pH is predominantly near neutral to slightly alkaline (pH 6.5–7.6). The Eh values vary between +594 and –444 mV, which suggests mildly oxidizing to moderate/strong reducing nature of groundwater environment in the BDP aquifers. The data plotted in an Eh–pH diagram (Fig. 3) indicates two clusters on the thermodynamic stability fields of trivalent As ($\text{H}_3\text{AsO}_3^\circ$ as unchanged species) as well as the field of pentavalent As (HAsO_4^{2-} as the protonated species). The groundwater types are generally Ca– HCO_3 or Ca–Mg– HCO_3 type, although Ca–Na– HCO_3 type and Na–Cl type water are also present locally in some areas (Bhattacharya et al. 1998b, 1999b; Ahmed 1999).

Major ion chemistry in groundwater is dominated by bicarbonate (320–600 mg L^{-1}) and shows an apparent depth and lithological control (Bhattacharya et al. 2001; Routh et al. 2000). Sulfate concentrations are by and large low ($\leq 3 \text{ mg L}^{-1}$) except for one location in Munshiganj district where a sulfate concentration was found to be high (8.5 mg L^{-1}). Nitrate concentrations are also low in the groundwater samples ($\leq 0.22 \text{ mg L}^{-1}$). On the other hand, phosphate concentrations (0.05–8.75 mg L^{-1}) are high in these analyzed groundwater samples.

Considerable variation in the concentration of total Fe (Fe_{tot}) is noted for the BDP groundwater of Bangladesh (0.4 to 15.7 mg L^{-1}). Total arsenic (As_{tot}) concentration in the analyzed wells varies between 2.5 and 846 $\mu\text{g L}^{-1}$. Arsenic(III) is the dominant species and accounts for about 67–99% of the total arsenic in most water samples. In certain wells, however, arsenic(V) dominates. Arsenic concentrations above the drinking water standards are concentrated at depths of <150 m. Deeper aquifers (>150 m) in general produce groundwater with arsenic concentrations below the WHO limit 0.01 mg L^{-1} (Bhattacharya and Jacks 2000); groundwater chemistry of our samples and their collecting sites are given in Table 3 and Fig. 4, respectively.

Table 3. Groundwater chemistry in selected tubewells from BDP aquifers of Bangladesh.

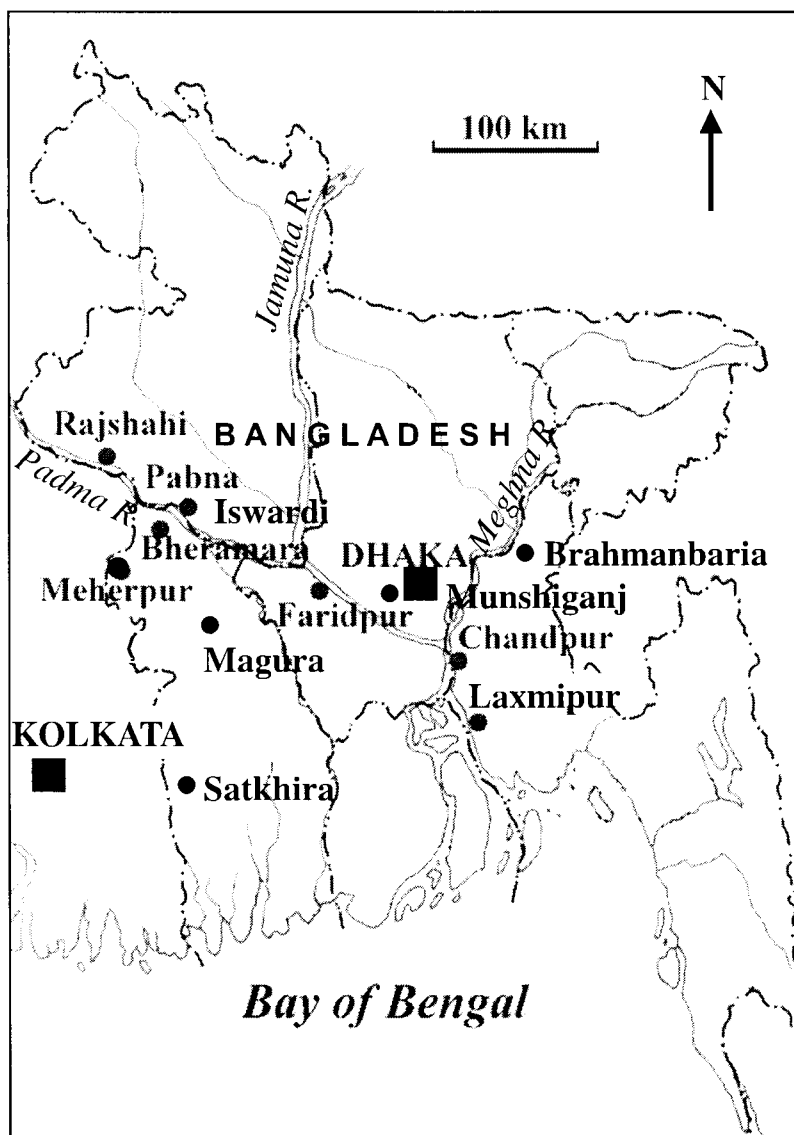
Location	Type	Depth (m)	T (°C)	pH	Eh (V)	EC ($\mu\text{S cm}^{-1}$)	HCO ₃ (mg L ⁻¹)	NO ₃ (mg L ⁻¹)	SO ₄ (mg L ⁻¹)	PO ₄ (mg L ⁻¹)	Fe _{tot} (mg L ⁻¹)	As _{tot} ($\mu\text{g L}^{-1}$)	As(III) ($\mu\text{g L}^{-1}$)	As(III) (%)	DOC (mg L ⁻¹)	References
Rajshahi																
Harian	Domestic	26.0	23.1	7.3	+0.342	738	392	0.10	bdl	2.30	9.7	152.4	133.2	87.4	1.61	This study
Harian	Domestic	28.0	24.9	6.8	+0.333	742	421	bdl	0.03	3.44	9.1	106.4	99.0	93.1	2.16	This study
Charghat	Domestic	36.6	26.6	7.0	+0.300	798	460	bdl	1.32	1.01	3.4	220.4	143.1	64.9	2.89	This study
Charghat	Domestic	45.0	25.8	7.0	+0.397	840	501	0.04	0.81	0.07	0.04	5.3	2.6	49.1	2.14	This study
Ishwardi																
CharRuppur	Domestic	28.5	22.7	7.1	-0.055	1312	739	bdl	0.12	0.69	3.3	529.6	476.0	89.9	2.07	This study
CharRuppur	Domestic	81.1	25.2	7.1	+0.194	760	445	bdl	0.76	0.08	0.6	4.38	3.78	86.3	2.88	This study
Meherpur																
Ujalpur	Domestic	28.5	25.8	7.2	-0.254	812	484	0.22	0.18	2.38	9.4	111.5	110.7	99.2	1.30	This study
Ujalpur	Irrigation	75.1	27.2	7.2	-0.444	767	462	bdl	0.02	1.32	2.8	96.9	88.1	90.9	1.73	This study
Faridpur																
Pourasabha	Domestic	21.0	25.2	7.6	-0.300	538	290	0.12	0.15	2.44	4.2	265.8	258.9	97.4	2.22	This study
Komarapur	Domestic	13.7	25.2	7.1	+0.157	640	449	0.07	2.83	0.18	0.15	22.3	nd	—	1.25	This study
Komarapur	Domestic	15.2	25.2	7.0	+0.594	630	379	0.04	1.77	0.54	1.48	54.2	51.0	94.1	1.56	This study
Komarapur	Domestic	21.3	26.0	7.0	+0.005	800	526	0.04	0.44	0.64	0.40	62.8	nd	—	3.06	This study
Komarapur	Domestic	228.6	26.2	7.1	+0.282	1180	697	0.05	bdl	0.27	0.55	110.0	100.4	91.3	7.29	This study
Dhuldi	Domestic	13.7	24.6	7.9	+0.157	860	611	0.06	0.04	2.63	15.7	845.6	812.0	96.0	4.14	This study
Rajapur																
Dhuldi	Domestic	15.2	25.1	7.2	+0.337	810	552	bdl	bdl	2.73	14.4	825.3	816.0	98.9	3.91	This study
Majhchar																
Rajapur	Domestic	16.8	25.2	7.1	-0.118	700	442	0.05	0.07	2.25	6.68	737.9	680.7	92.3	3.09	This study
Kourpur																
Majhchar	Domestic	21.3	24.8	7.1	-0.019	630	475	0.12	0.17	0.63	4.42	22.3	21.4	96.0	2.01	This study
Kourpur	Domestic	243.8	26.1	6.5	-0.124	1970	209	0.78	8.52	0.35	1.20	10.2	2.6	25.4	1.15	This study
Munshiganj																
Srinagar	Domestic	54.9	25.7	6.9	-0.326	620	345	0.04	0.01	6.35	3.19	147.7	144.8	98.1	3.53	This study
Maowa	Domestic	67.1	25.6	6.4	+0.104	1300	874	0.13	0.11	10.43	5.85	197.3	187.5	94.0	14.2	This study
Srinagar	Domestic	243.8	26.1	6.5	-0.124	1970	209	0.78	8.52	0.35	1.20	10.2	2.6	25.4	1.15	This study

Table 3. (concluded).

Location	Type	Depth (m)	T (°C)	pH	Eh (V)	EC ($\mu\text{S cm}^{-1}$)	HCO ₃ (mg L ⁻¹)	NO ₃ (mg L ⁻¹)	SO ₄ (mg L ⁻¹)	PO ₄ (mg L ⁻¹)	Fe _{tot} (mg L ⁻¹)	As _{tot} ($\mu\text{g L}^{-1}$)	As(III) ($\mu\text{g L}^{-1}$)	As(III) (%)	DOC (mg L ⁻¹)	References
Magura																
Pourasabha	Domestic	33.5	25.3	6.9	-0.106	740	487	0.03	0.20	1.48	5.53	53.5	47.3	88.4	1.82	This study
Pourasabha	Domestic	45.7	25.2	7.0	+0.202	710	476	bdl	bdl	2.16	3.60	40.2	nd	—	1.77	This study
Brahman Baria																
Sonarampur	Domestic	36.6	25.2	7.2	+0.137	520	275	bdl	0.202	15.3	2.04	327.8	320.9	97.9	8.13	This study
Sonarampur	Domestic	22.9	24.2	7.3	+0.182	510	297	0.12	0.05	8.75	2.28	234.6	226.7	96.6	3.05	This study
Kuttapara, Sadar	Domestic	51.8	24.9	7.1	+0.309	620	314	bdl	bdl	2.16	4.13	207.5	196.2	94.6	4.52	This study
Haldarpara, Sadar	Domestic	97.5	24.9	7.2	+0.118	550	235	0.04	0.06	0.05	0.01	2.25	nd	—	1.30	This study
Astagram, Sadar	Domestic	51.8	25.4	7.3	+0.038	410	263	0.06	0.04	3.87	0.56	186.6	nd	—	3.77	This study
Medda, Sadar	Domestic	73.2	26.5	7.1	+0.153	430	261	0.03	0.07	5.47	2.90	99.9	nd	—	3.52	This study
Hathata, Sadar	Domestic	42.7	26.2	7.1	-0.118	650	404	bdl	0.02	4.06	4.67	372.7	nd	—	6.61	This study
Saikhira																
Tala	Domestic	25.9	27.4	6.7	-0.010	611	377	0.05	bdl	5.84	3.97	136.6	105.6	77.3	2.14	Mazumdar 2000
Tala	Domestic	29.0	27.1	6.7	+0.013	941	554	0.21	bdl	5.94	10.7	182.9	134.8	73.7	4.34	Mazumdar 2000
Tala	Domestic	54.9	27.7	6.8	-0.008	752	423	0.04	bdl	5.70	3.6	146.3	98.2	67.1	2.87	Mazumdar 2000
Tala	Domestic	62.5	27.6	6.9	+0.003	900	567	0.07	0.07	4.09	5.4	170.9	125.2	73.2	3.43	Mazumdar 2000
Laxmipur																
Shamsherabad	Domestic	7.9	25.3	7.1	+0.304	739	362	bdl	0.051	1.81	3.38	174.9	165.1	94.4	1.87	This study
Hazirpara	Public	11.7	25.8	7.0	-0.271	640	343	0.02	0.04	4.79	2.97	210.5	153.0	72.7	2.03	This study
Hazirpara	Domestic	255.3	26.2	6.7	-0.107	307	207	bdl	bdl	0.75	1.84	5.14	0.66	12.9	3.27	This study

Notes: EC, electrical conductivity; bdl, below detection limit; nd, not detected.

Fig. 4. Groundwater sampling sites (Rajshahi, Iswardi, Faridpur, Meherpur, Munshiganj, Magura, Brahman Baria, Satkhira, Laxmipur) in Bangladesh (this study).



Fe_{tot} shows fairly high coefficients of linear regression with HCO_3 ($r^2 = 0.57$, Fig. 5a) as well as with PO_4 ($r^2 = 0.50$; $p < 0.001$, Fig. 5b). The linear regression coefficient is comparatively low between Fe_{tot} and As_{tot} ($r^2 = 0.42$; $p < 0.001$, Fig. 5d) in the groundwater samples. High HCO_3 concentrations are fairly well correlated with the levels of dissolved organic carbon (DOC) (1.15 – 14.2 $mg\ L^{-1}$) in groundwater ($p < 0.05$; $r^2 = 0.38$, Fig. 5c). $As(III)$ and Fe_{tot} concentrations in the groundwater show two distinct trends of variations with DOC (Fig. 6 a and b). The source of DOC is however not well known and needs to be investigated further, but the sedimentary organic matter (OM) in the aquifers may be thought to be active source for the DOC in the BDP groundwaters.

In a reducing environment and with low As:Fe ratios, increase in arsenic concentration in groundwater is related to the microbial reduction of arsenate, $As(V)$, to arsenite, $As(III)$ (Langner and Inskeep

Fig. 5. Chemical characteristics of groundwater from BDP aquifers in Bangladesh ($n = 36$) showing the relation between (a) Fe_{tot} and HCO_3 , (b) PO_4 and Fe_{tot} , (c) dissolved organic carbon and alkalinity, and (d) Fe_{tot} and As_{tot} .

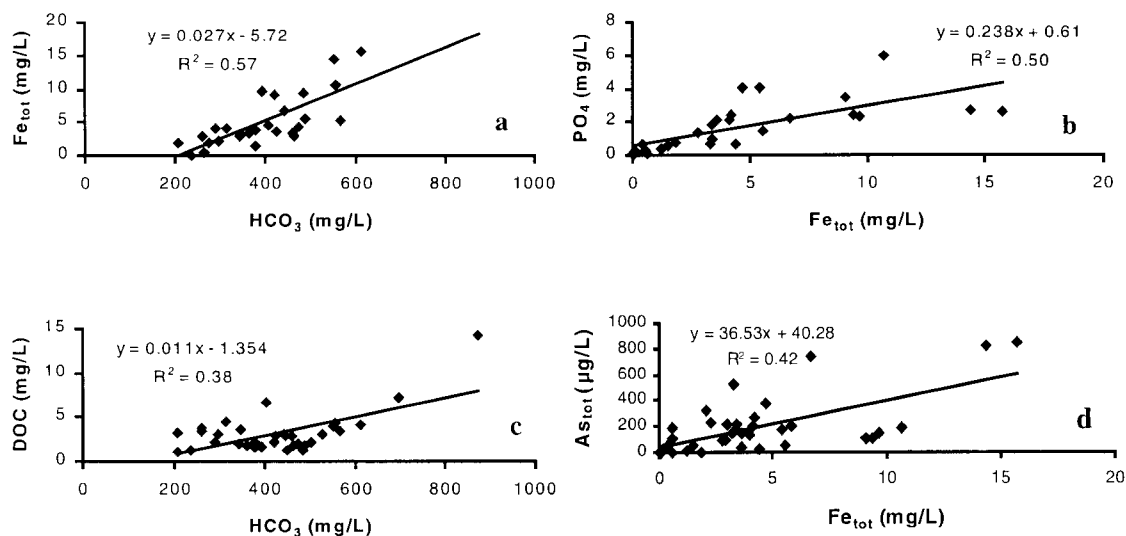
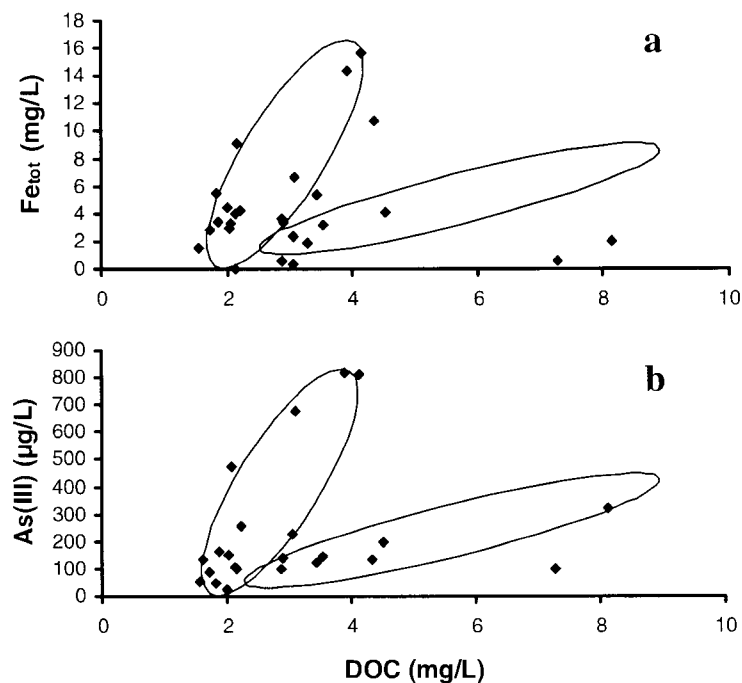


Fig. 6. Sympathetic variation of Fe_{tot} and As(III) with dissolved organic carbon in BDP groundwaters from Bangladesh. The data points with distinct trends suggest that both As(III) and Fe_{tot} are mobilized from the aquifers by DOC consequent to the microbial reduction of the sedimentary organic matter.



2000), and reductive dissolution of Fe(III) species, which results in the release of arsenic and other trace elements as well as phosphate in groundwater. High DOC levels and dominance of As(III) species in near-neutral and anoxic BDP groundwaters also suggest that reduction of OM by microorganisms may facilitate biomethylation processes, which may result in the conversion of As(V) to the more soluble As(III) species, thereby affecting the mobility and translocation of arsenic (Routh et al. 2000).

4.3.2. Nature and chemistry of the alluvial sediments in the Bengal Delta Plain

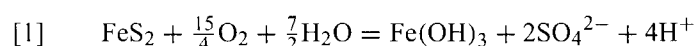
Geochemical analysis of the alluvial aquifers sediments from the BDP have revealed high concentrations of arsenic elevated by factors of 2.5 to >70 as compared to the average continental crust. Total arsenic (HNO_3 extraction, As_{NO_3}) concentrations in the sediments have been estimated to be in the range of 5–14 mg kg^{-1} at different locations and depths. Some of the aquifer sediment from Tala (in Satkhira district) revealed high concentrations of arsenic (294 mg kg^{-1}). Considerable variations are also noted for the distribution of Fe_{NO_3} (57.8–334.9 g kg^{-1}), Al_{NO_3} (36.9–223 g kg^{-1}), and P_{NO_3} (1.5–5.9 g kg^{-1}).

Significant positive Spearman rank correlations are noted amongst As_{NO_3} and Fe_{NO_3} ($p < 0.001$), Al_{NO_3} ($p < 0.05$), Mn_{NO_3} ($p < 0.001$), and P_{NO_3} ($p < 0.05$) in the aquifer sediments (Fig. 7 *a–d*). The dominance of Fe-oxyhydroxides in the sediments is revealed by higher quantities of oxalate extractable iron ($\text{Fe}_{\text{ox}} = 4.2\text{--}174 \text{ g kg}^{-1}$) than aluminum ($\text{Al}_{\text{ox}} = 0.6\text{--}43.7 \text{ g kg}^{-1}$) or manganese ($\text{Mn}_{\text{ox}} = 0.1\text{--}4.5 \text{ g kg}^{-1}$). The amount of oxalate extraction arsenic (As_{ox}) varies between 1.7 and 86.4 mg kg^{-1} . Strong correlation (Fig. 8 *a* and *b*) is also noted amongst Fe_{NO_3} and Fe_{ox} ($p < 0.001$) and As_{NO_3} and As_{ox} ($p < 0.001$). Among the oxalate extractable fractions, high correlation (Fig. 9 *a–d*) exist between As_{ox} and Fe_{ox} ($p < 0.01$), Al_{ox} ($p < 0.05$), Mn_{ox} ($p < 0.001$), and P_{ox} ($p < 0.05$). Correlation between oxalate extractable arsenic and the secondary amorphous Fe-, Al- and Mn-oxyhydroxides in the sediments, however, implies that they act as the principal adsorbent for arsenic in the BDP aquifers sediments (Bhattacharya et al. 2001).

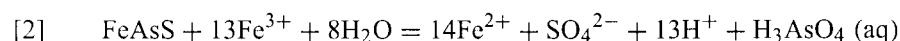
4.3.3. Mechanisms of arsenic mobilization

Though the causes of widespread occurrences of arsenic in groundwater of the BDP are a matter of debate, the basic geochemical processes that might be responsible for the problem are well established (von Brömssen 1999; Bhattacharya et al. 1999*a*, 1999*b*, 2001; Bhattacharya and Mukherjee 2000; Mazumdar 2000; Nickson et al. 2000).

Pyrite oxidation is cited as a plausible hypothesis for the widespread arsenic contamination of the BDP groundwater (Das et al. 1996; Roy Chowdhury et al. 1999 and many others). Pyrite is oxidized in aquifer sediments where groundwater is oxygenated according to the reaction



while the oxidation of arsenopyrite follows the reaction



Both these reactions produce water with high concentrations of Fe^{2+} and SO_4^{2-} accompanied by a decrease in pH. Validity of this hypothesis is under sharp criticism because the groundwaters in this region are (i) anoxic, with no or negligible dissolved oxygen; (ii) at near neutral pH; (iii) strongly alkaline; (iv) characterized by low sulfate concentrations with no relationship with arsenic; and (v) characterized by virtual absence of nitrate (Bhattacharya et al. 1999*b*; BGS/MML 1999*a*).

Mobilization of arsenic from the alluvial aquifers is primarily due to the mechanism of desorption of arsenic oxyanions from the surface of Fe-oxyhydroxides as well as reductive dissolution of the Fe-oxyhydroxides in the sediments (Bhattacharya et al. 1996, 1997; Nickson et al. 1998; BGS/MML 1999*a*). The present study suggests that the bulk of the arsenic in the sediments is quantitatively related to the amounts of the amorphous Fe-, Al-, and Mn-oxyhydroxides in the sediments. These surface reactive compounds are characterized by a positive charge and therefore act as a strong adsorbent for the arsenic oxyanions (Pierce and Moore 1982). Increased pH of groundwater releases arsenic oxyanions by desorption, and the inflow of DOC-charged, anoxic water from the cultivated wetlands (rice fields) facilitates the dissolution of Fe-oxyhydroxides under the prevalent reducing conditions.

Fig. 7. Relationship between the HNO₃ extractable fraction of As (As_{NO₃}) and (a) Fe_{NO₃}, (b) Al_{NO₃}, (c) Mn_{NO₃}, and (d) P_{NO₃} in the alluvial aquifer sediments from Chapai Nawabganj and Satkhira districts of Bangladesh.

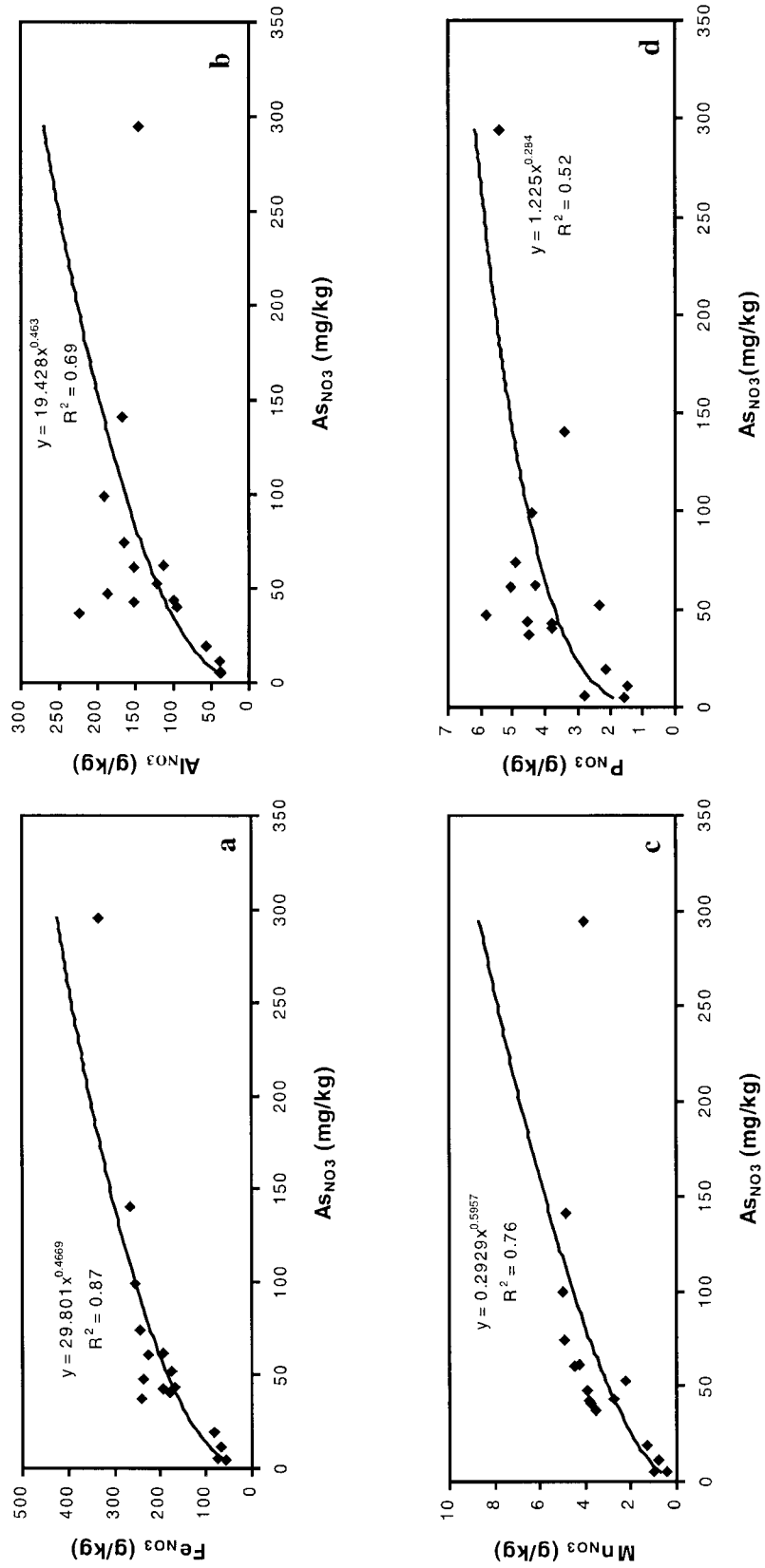
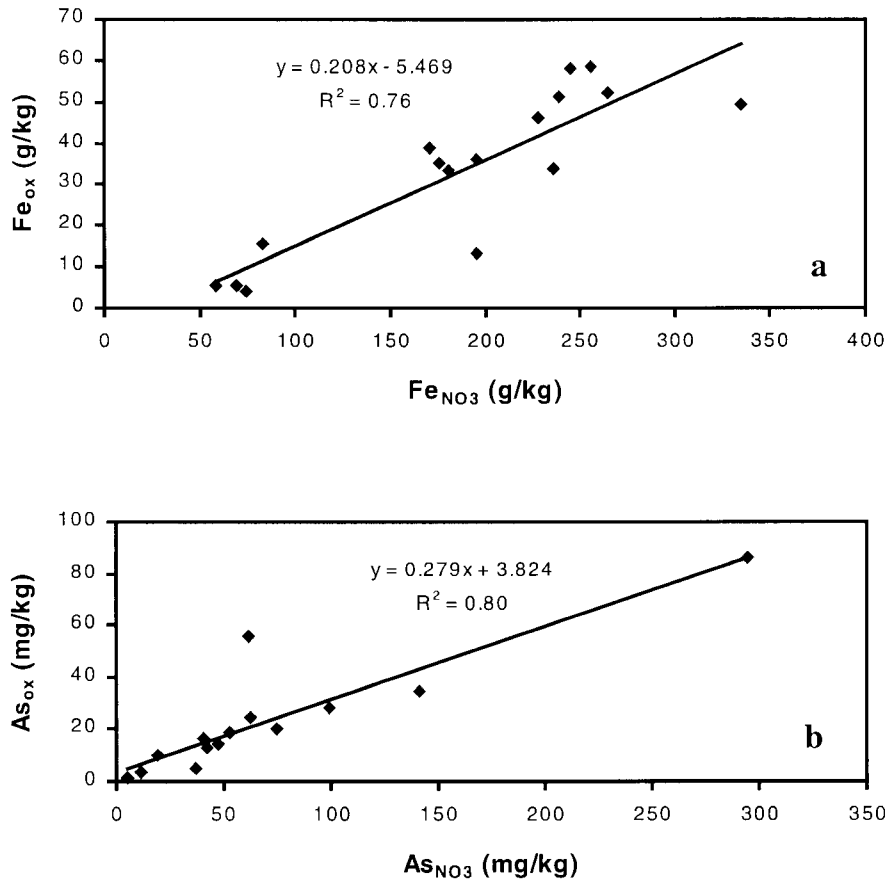
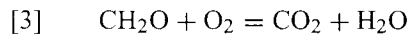


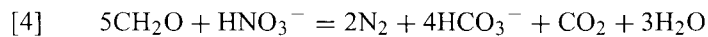
Fig. 8. Relation between the HNO₃ extractable and oxlate extractable fraction of Fe and As in the aquifer sediments.

Consumption of the dissolved oxygen and other oxidants (such as nitrate) mediate the oxidation of the sedimentary OM in the aquifers, thereby increasing the groundwater alkalinity (Bhattacharya et al. 1997, 2001). High alkaline groundwater in the BDP aquifers may thus be produced by (i) oxidation of organic matter in the presence of oxygen, expressed by the reaction



and

(ii) bacterially mediated reduction of nitrate by organic matter, following the reaction



Bacterial reduction plays an important role in these aquifers because both nitrate and sulfate concentrations are low. High NH₄⁺ concentrations in these groundwaters (Mazumdar 2000; Bhattacharya and Jacks 2000) also tend to suggest that bacterially mediated nitrate reduction processes are taking place in these aquifers (for example, Zehnder 1988; Smith et al. 1991). Studies by Larsson and Leiss (1997) and Bhattacharya et al. (1997) also pointed out that sedimentary arsenic can be mobilized by microbial degradation in the presence of organic substrates in reducing aquifers. Our studies reveal well-defined correlation between the concentrations of bicarbonate with Fe_{tot} as well as the DOC. Sedimentary and dissolved OM serve as electron donors that control the key redox reactions (e.g., NO₃⁻ and SO₄²⁻ reduction, dissolution of Fe(OH)₃) in the aquifers. The correlation amongst Fe_{tot} and As_{tot}, as well as

Fig. 9. Relationship between the oxalate extractable As (As_{ox}) and (a) amorphous Fe-oxyhydroxides (Fe_{ox}), (b) Al-oxyhydroxides (Al_{ox}), (c) amorphous Mn-oxides (Mn_{ox}), and (d) oxalate extracted P (P_{ox}) in the aquifer sediments.

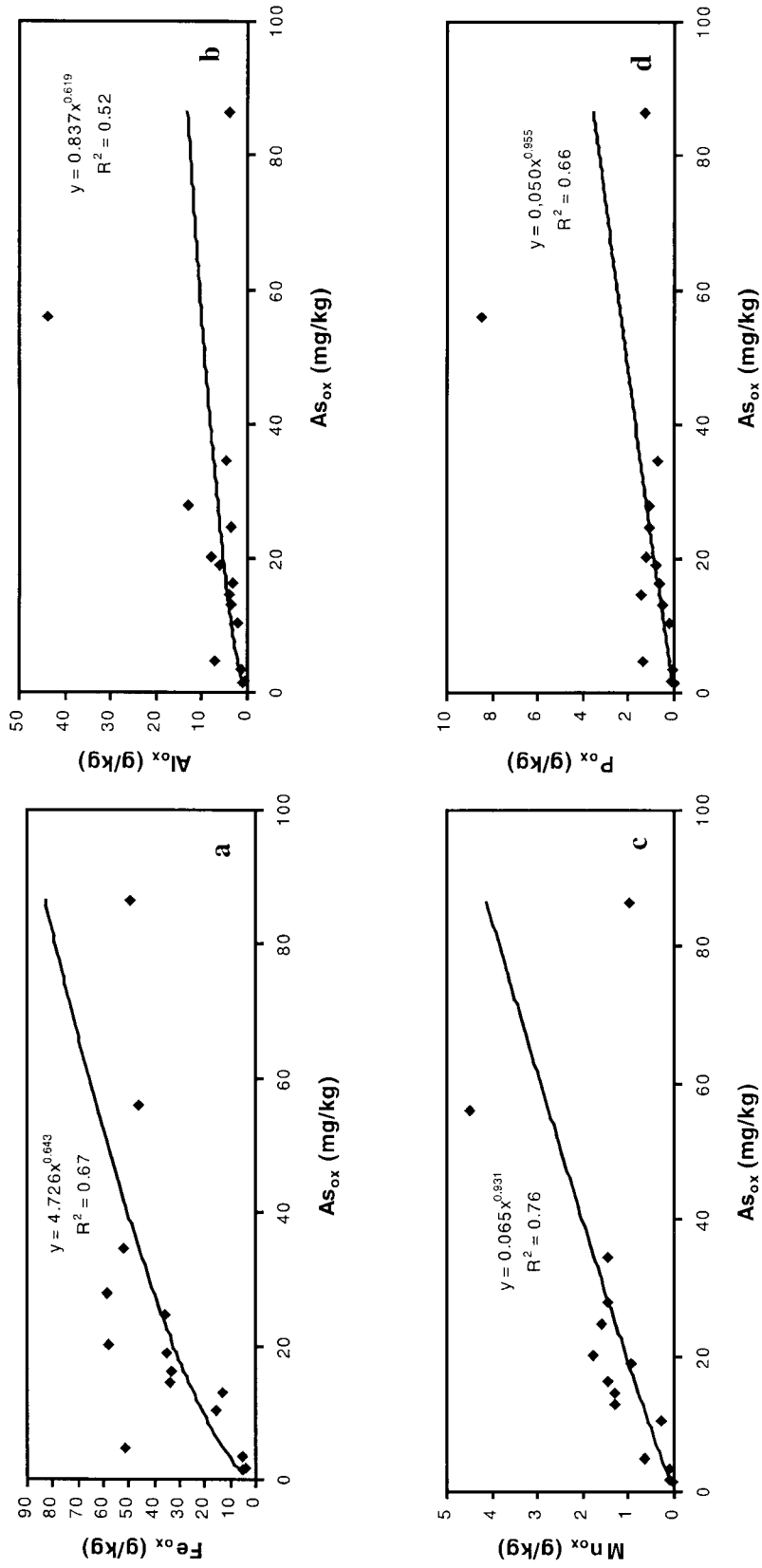


Table 4. Estimated daily intake of arsenic in the affected areas of Bangladesh.

Population at risk of the affected districts ^a	75 million
Drinking water habits including rice water (L d ⁻¹) ^b	5
As concentration ($\mu\text{g L}^{-1}$)	>50–1000
As intake per person per day ($\mu\text{g d}^{-1}$ person ⁻¹)	>250–5000
Intake for affected population (ca. 75 million) (t a ⁻¹)	5.48–11

^aDainichi Consultant, Inc. (1999).^bBGS/MML (1999a).

PO₄ in the groundwater, suggest that both, as well as PO₄³⁻, are released to the groundwater following reductive dissolution of Fe(OH)₃. However, the correlation among these parameters can be masked because of the precipitation of siderite (FeCO₃) and to some extent vivianite (Fe(PO₄)₂·8H₂O), which act as the sink for Fe²⁺ in the reducing aquifers (Sracek et al. 2000). Arsenic-rich pyrites (framboidal pyrite) on the other hand are formed diagenetically in the clayey sediments in the sulfate reducing zone. These secondary sulfides, however, act as a sink for arsenic and remain stable under reducing conditions (Ahmed 1999; Bhattacharya et al. 1999b). All this evidence suggests that arsenic mobilization from the alluvial aquifers in the BDP occurs predominantly under the reducing environment and the mechanism is controlled by the dissolution of Fe-, Al-, and (or) Mn-oxyhydroxides due to the biogeochemical interactions in the aquifers.

5. The role of arsenic in biology, health, and society of Bangladesh

5.1. Drinking water

For the last 2000 years, arsenic has been used as a medicine while at the same time being known to be a poison. In the last decades, high levels of arsenic concentration in groundwater from natural and industrial sources have created environmental pressures in different regions of the globe. In many areas, groundwater is used for drinking purpose, as it is safe. Generally, drinking water and the food chain are the main pathways by which one can be exposed to the poisonous effects of arsenic. It has already been discussed how the arsenic level in groundwater in Bangladesh varies from one region to another because of geological and geomorphological nature of the areas as well as the fact that the aquifers are shallow (see Table 3 and Fig.1). The information indicates how groundwater arsenic concentration varies with shallow aquifers (<150 m). Even old and new types of wells generate different results, which cannot be explained. However, different studies indicate that in the affected areas, groundwater arsenic concentration varies between >50 and 1000 $\mu\text{g L}^{-1}$ (maximum permissible limit in Bangladesh: 50 $\mu\text{g L}^{-1}$). Occasionally, very high concentrations of arsenic in shallow tubewells in Pabna (14 mg As L⁻¹) and in Kushtia (9 mg As L⁻¹) were reported (The New Nation 1996; DCH 1997). It is possible to estimate the daily intake of arsenic from drinking water in the affected areas and this is shown in Table 4. Daily inorganic arsenic intake for the adult population in the affected areas varies between >250 and 5000 $\mu\text{g day}^{-1}$ as compared to 5 $\mu\text{g day}^{-1}$ in the United States (US EPA 1982). This high intake originates not only from drinking water, but also from drinking rice-water and water used for cooking.

5.2. Food chain

Use of high arsenic content groundwater for irrigation has resulted in the accumulation of arsenic in crops and plants. Based on the studies carried out so far, it is significant to note that high concentrations of arsenic in vegetables and rice in affected areas, such as Chandipur village in Laximpur (Table 5), indicate that dietary habits are also responsible for arsenic problems for the people of Bangladesh (Huq et al. 2001). The arsenic concentration values of vegetables grown in Bangladesh should not be

Table 5. Arsenic concentration ($\mu\text{g g}^{-1}$) in vegetables in an affected area in Bangladesh (Huq et al. 2001). Figures rounded.

Samples	Arsenic concentration ($\mu\text{g g}^{-1}$)		
	Min	Max	Average
Rice (<i>Oryza sativa</i>)	0.44	0.5	0.47
Kochu, leaf (<i>Colocasia antiquorum</i>)	0.11	0.11	0.11
Kochu, stem (<i>Colocasia antiquorum</i>)	0.74	0.92	0.83
Kochu, creeper (<i>Colocasia antiquorum</i>)	0.21	0.23	0.22
Gourd, leaf (<i>Lagenoria siceraria</i>)	1.88	2.01	1.94
Brinjal (<i>Solanum melongena</i>)	0.18	0.20	0.19
Sweet potato, leaf (<i>Ipomea batatus</i>)	0.38	0.41	0.40
Pui shak (<i>Basilia alba</i>)	0.30	0.33	0.31
Kalmi shak (<i>Ipomea aquatica</i>)	0.29	0.35	0.32
Papaya (<i>Carica papaya</i>)	0.19	0.26	0.22
Red pumpkin (<i>Cucurbita maxima</i>)	0.20	0.27	0.24

compared to that of vegetables grown in Europe, as the species grown in Bangladesh are completely different to those grown in Europe.

Fish constitutes an important part of the diet in Bangladesh. Concentration of arsenic in water is not always responsible for the arsenic level in fish species. Most important is how a metal is transferred tropically through the food webs and, for example, whether it is biomagnifiable or biodiminshable. Fish species have different food habits. Planktivorous species (e.g., alewives and killifish) may contain considerably more arsenic than omnivorous species (bluegill and yellow perch) or piscivorous species (large mouth bass) (Fig. 10) (Chen and Folt 2000). Although there are no values for the arsenic concentrations in fish in Bangladesh, it is important to note that much of the arsenic present in fish and shellfish is in either the fat-soluble or water-soluble organoarsenic form that is essentially non-toxic (Lunde 1973). A recent study in the Netherlands suggests that in seafood 0.1–41% of the arsenic is present as inorganic arsenic (Vaessen and van Ooik 1989; ATSDR 1993). Canadian studies have indicated that the arsenic in sea fish was 74–91% organic arsenic, whereas in lake fish this was 71–85% (Lawrence et al. 1986). In Canadian ocean fish and lake fish, total arsenic was reported at 4.2 and 0.08 mg kg^{-1} , respectively (Lawrence et al. 1986), whereas for Finland the total arsenic concentration was between 0.61 and 0.12 mg kg^{-1} (Liukkonen-Lilja 1993). Hence, we believe that also in Bangladesh total arsenic concentration in river or ocean fish will vary from one species to another. Total arsenic (inorganic and organic) intake derived from foods for a number countries are given in Table 6. In Japan, higher values have been cited; these are due to presence of a higher percentage of organic arsenic in fish species.

5.3. Impact of high arsenic ingestion

The toxic properties of arsenic have been known for centuries. The symptoms of acute intoxication of arsenic have been reported in many countries, including Argentina, Bolivia, Chile, Taiwan, India, Bangladesh, and others (Borgoño et al. 1977; Guha Mazumder et al. 1988; 1992). According to FAO/WHO (1998), the daily acceptable maximum range of arsenic intake for a healthy 70 kg person via air, water, and food is as follows: air, 0.8–30 ng m^{-3} ; water, 10–50 $\mu\text{g L}^{-1}$; and food, 0.8–120 $\mu\text{g kg}^{-1}$. An interesting study has been done by Tondel et al. (1999) regarding arsenic associated skin lesions from four villages in the districts Faridpur, Jessore, Narayanganj, and Nawabganj in Bangladesh. Their study indicated high prevalence rate of hyperpigmentation for males (22.7/100) and females (11.5/100) who drink tubewell water containing arsenic of concentrations $>800 \mu\text{g L}^{-1}$. In these areas, maximum arsenic concentration in drinking water was 3400 $\mu\text{g L}^{-1}$. Analyses of arsenic in urine (0.05–9.42 mg L^{-1}), hair (1.1–19.84 mg kg^{-1}), fingernails (1.3–33.98 mg kg^{-1}), and skin

Fig. 10. Variation of arsenic concentration in planktivores (AWF, alewife; KLF, Killfish), omnivores (BLCR, Black crappie; BGS, Bluegill sunfish; YP Yellow perch), and piscivores (LMB, Largemouth bass) fish species. Fish species are from lakes in New Hampshire, U.S.A. (modified from Chen and Folt 2000).

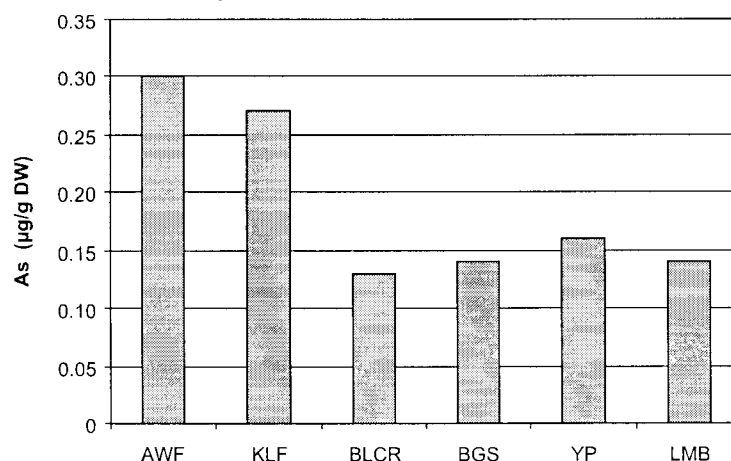


Table 6. Total arsenic (inorganic + organic) intake from foods in selected countries. Source: Liukkonen-Lilja (1993).

Country	Arsenic intake ($\mu\text{g As d}^{-1}$)	Year
Switzerland	30	1987
Holland	38	1989
Canada	49	1993
Finland	58	1980
Germany (West)	67	1983
England	89	1982
Denmark	118	1990
Japan	197	1990
PTWI ^a	130 ^b	

^aPTWI, Provisional Tolerable Weekly Intake.

^bInorganic As intake per day is based on the weight of a person of 60 kg.

scale ($0.6\text{--}53.39 \text{ mg kg}^{-1}$) indicate severe contamination among the affected population in Bangladesh (Chowdhury et al. 1999; Karim 2000). The accepted safe limits include urine, $0.005\text{--}0.04 \text{ mg d}^{-1}$; hair, $0.08\text{--}0.25 \text{ mg kg}^{-1}$; and fingernails, $0.43\text{--}1.08 \text{ mg kg}^{-1}$ (Arnold et al. 1990; Samanta et al. 1999; Karim 2000). Dose response relationships between inorganic arsenic exposure from tubewell water and risk of hypertension, diabetes mellitus, and keratosis were established in Bangladesh and many other countries, including Taiwan and Sweden (Lai et al. 1994; Rahman et al. 1996, 1998, 1999). At least 7000 people are suffering from arsenicosis and 30–70 million people are thought to be exposed to toxicity by arsenic (DCH 1998; Dainichi Consultant, Inc. 1999). The toxic effects of inorganic arsenic are skin lesions, hyperkeratosis, skin cancer, liver disease, and hyperpigmentation (WHO 1981; Morton and Dunnette 1994). High concentration of arsenic ($1,860 \mu\text{g L}^{-1}$) was also reported at the Zhi Ji Liang village of Inner Mongolia, China. In three villages of that area more than 168 patients were diagnosed as suffering from chronic arsenic poisoning (e.g., hyperkeratosis, hyperpigmentation, and skin lesions) (Luo et al. 1997). Toxicity is determined on the basis of ingestion of arsenic compounds and their excretion from the body. It is expected that 40–60% of the arsenic is retained by the human body (Farmer and Johnson

1990). The half life of arsenic in humans is cited at 28 to 49 days (Pomroy et al. 1980; Buchet et al. 1980).

5.4. Social implications

Arsenic, king of poisons, has brought deep social problems to the communities of Bangladesh. The country is highly populated, which has forced the nation to intense use of land, forest, fisheries, and water resources. This has created severe environmental pressure on cultivation, soil fertility, excess extraction of water for irrigation, commercial fish farming, and use of agricultural chemicals. It is obvious that national economics are based on the commodities and services derived from ecosystems. But little attention has been paid to the ecosystem. Pressure is mounting quickly because of the disturbed socioeconomic conditions of the country. Some people mistake skin disorders resulting from arsenocosis for leprosy. Married women affected by arsenic are sent back to their parents with their children. Affected children are prevented from attending school, and many young men are deprived of jobs when employers find black spots on their body and hands. Affected young unmarried women and men are advised to stay unmarried. They are not allowed even to go to their neighbors to fetch water. Social problems in the affected areas know no bounds (Biswas et al. 1998; Samanta et al. 1999). It is observed that poor people are most badly affected by arsenic concentration in groundwater. But still much needs to be understood regarding the impact of arsenic poisoning. Some family members or communities are affected from arsenic poisoning, whereas members in the same family or community who are subject to the same contamination are not. Campaigns are currently ongoing to make people understand that those sick from arsenic are not a threat to others, but deserve attention and care. UNICEF has launched the Facts for Life (FfL) program to promote "awareness about the need for sensitivity to emerging social issues, such as children and women being ostracized when they show signs of being affected by arsenic" (Anon by Bangladesh-UNICEF).

6. Arsenic removal

Arsenic removal technologies are available, but cost effective methods are necessary for the people of Bangladesh. As mentioned above, arsenic is present in the aquatic environment as reduced As(III) and oxidized As(V). As the chemical forms of arsenic are different, it is necessary to estimate the speciation of arsenic before adopting any removal technology. In Bangladesh, groundwater contains 43–98% inorganic arsenic as As(III) (Rasul et al. 1999), which is more toxic than As(V) (Korte and Fernando 1991). It has been reported that it is not possible to remove arsenic(III) from the water in question, but it must first be oxidized to arsenic(V) (Gottschalk et al. 1992). Figure 11 summarizes the available arsenic removal methods. It is worth mentioning that UNICEF and DPHE (Department of Public Health Engineering in Bangladesh) have financed a project at the Bangladesh University of Engineering and Technology to evaluate the use of activated alumina for arsenic removal. In addition, groundwater arsenic research is ongoing in the Division of Land and Water Resources at the Royal Institute of Technology, Stockholm, Sweden. In this institution, low-cost techniques for removal of arsenic from groundwater have been identified. These include (i) auto-attenuation, (ii) use of geological materials as natural adsorbent for arsenic, such as laterite or Fe-rich oxisols, and (iii) re-infiltration of groundwater to the aquifers following aeration. Studies by Larsson et al. (1999) indicated that laterite (consisting mainly of hydrous oxides of Fe and Al and minor amounts of Mn and Ti) adsorbed 50–90% of arsenic from contaminated groundwater (Fig. 12). Among the above-mentioned techniques, the most encouraging method to purify arsenic-laden groundwater is auto-attenuation. Details of these processes may be found elsewhere (Larsson et al. 1999; Bhattacharya et al. 2000).

The United States Environmental Protection Agency (USEPA) has recently identified three promising methods for the removal of arsenic from groundwater: (i) ion exchange with brine recycles, (ii) attenuation method, and (iii) coagulation/filtration in the presence of iron.

Fig. 11. Various groundwater arsenic removal principles (BGS/MML 1999a).

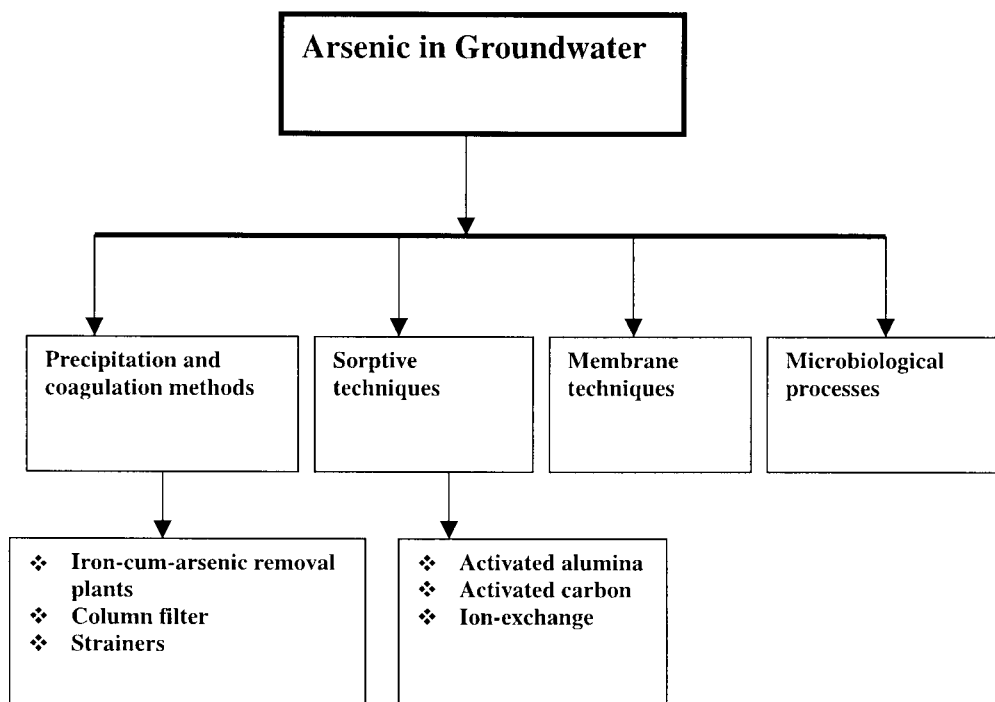
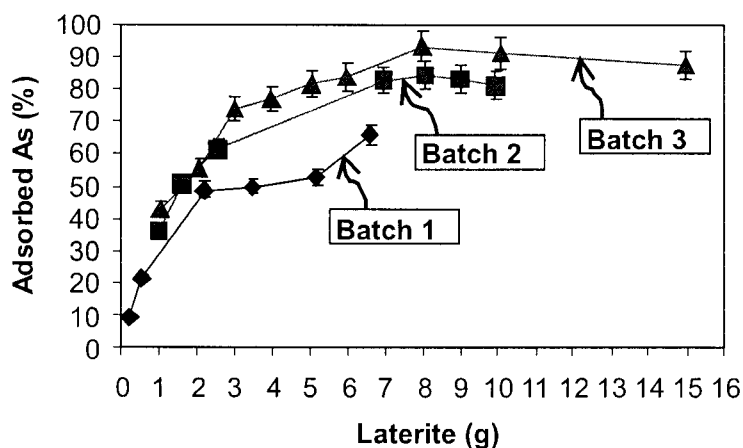


Fig. 12. Amount of arsenic adsorbed on laterite (in percent) from contaminated groundwaters of BDP (modified from Larsson et al. 1997). Results shown above are based on three batch tests where 100 mL contaminated groundwater ($As_{tot} = 334 \mu\text{g L}^{-1}$) was treated with different amounts of laterite. The bars ($\pm 5\%$) represent the range of variability for arsenic adsorption in the batch tests.



On the other hand, a group of engineers from the Bengal Engineering College, Howrah, West Bengal, India, have developed a unit completely manually operated and not requiring electricity for handwells and hand pumps. Arsenic removal columns are installed at the tubewells using activated alumina. Once every 4 months the regeneration of the unit with caustic soda is necessary. Each unit costs about US\$1250 and can supply arsenic free water for 200 to 300 households. Lifetime of the units, without any maintenance, is about 10 years, and the arsenic-containing sludge can be stored in a pit (121 × 121 × 121 cm) for minimal 8 to 10 years (Anon 2000b).

Table 7. Concentration of metals in test run groundwater samples from Bangladesh before and after filtration by 3-pitcher filtration method (Khan et al. 2000)

Elements	Before filtration (mg L ⁻¹)	After filtration (mg L ⁻¹)
Aluminum	0.031 ± 0.022	0.023 ± 0.011
Antimony	<0.013	<0.013
Arsenic	0.130–1.16	<0.016
Barium	0.166 ± 0.006	0.063 ± 0.01
Beryllium	<0.001	<0.001
Cadmium	<0.001	<0.001
Calcium	113 ± 2.5	59.1 ± 7.5
Chromium	<0.002	<0.002
Cobalt	<0.002	<0.002
Copper	0.008 ± 0.001	0.005 ± 0.002
Iron	6.897 ± 0.110	0.084 ± 0.002
Lead	0.005	0.006 ± 0.002
Magnesium	21.67 ± 0.55	23.17 ± 3.29
Manganese	0.723 ± 0.017	<0.001
Molybdenum	0.002 ± 0.001	0.003 ± 0.001
Nickel	<0.002	<0.002
Potassium	2.23 ± 0.23	6.71 ± 1.31
Selenium	<0.012	<0.012
Silver	<0.002	<0.002
Sodium	19.5 ± 1.14	26.7 ± 3.15
Strontium	0.287 ± 0.01	0.164 ± 0.02
Thallium	<0.067	<0.067
Tin	0.006 ± 0.004	0.011 ± 0.008
Vanadium	<0.001	0.006 ± 0.003
Zinc	0.08 ± 0.11	0.009 ± 0.002

Besides the above-mentioned developments, Khan et al. (2000) claimed that it is possible to reduce total inorganic arsenic and As(III) concentration below the detection limit of 2 and 10 $\mu\text{g L}^{-1}$, respectively, by using simple local ingredients such as coarse and fine sand, wood charcoal, and iron chips. The whole process is based on filtration and it is known as the “3-pitcher (locally known as 3-*kalshi*) water purification method”. Table 7 indicates the water quality after using this purification method. The authors believe that this is a cost-effective filtration method for removal of arsenic from contaminated groundwater. The process can be further developed by increasing the flow rate of water, but 6.2 L h⁻¹ is sufficient for a middle size family of 4–5 persons.

7. Conclusions

For thousand of years, groundwater has served as a unique and reliable source of potable water in developed as well as developing countries. In spite of enormous resources of water from surface and sizable precipitation in Bangladesh, the country has faced tremendous shortage of drinking water due to the risks of microbial contamination in surface water. Exploitation of groundwater resources started in Bangladesh during the 1960s and has increased dramatically in the past three decades to reach a goal for providing safe drinking water to nearly 97% of the rural population. Moreover, use of groundwater was also promoted to support wetland cultivation in the region. In 1993, arsenic was detected in groundwater at elevated concentrations ($>50 \mu\text{g L}^{-1}$), which has posed a serious threat to public health. Use of arsenic-contaminated groundwater in irrigation has, moreover, resulted in bioaccumulation of arsenic in crops and vegetables grown in the region.

Inorganic arsenic enters the ecosystem by natural and anthropogenic sources. Data on industrial emissions of arsenic to the atmosphere in Bangladesh are scarce, although a high concentration of arsenic (5000 mg kg^{-1}) has been reported in the soil in the vicinity of some industrial facilities. However, causes of groundwater arsenic contamination arising from anthropogenic sources is not applicable for such a wide-scale arsenic problem in the Holocene aquifers of the BDP. Arsenic concentration in groundwater of the BDP is by and large prevalent from the Holocene sedimentary aquifers. Wells placed at the shallow aquifers ($>10\text{--}150 \text{ m}$) mostly yield water with high arsenic concentration, but the concentration of arsenic is found to decrease with depth. This observation is, however, quite inadequate for generalization, as factors controlling the mobilization of arsenic at the sediment–water interface are dependent on the lithological and biogeochemical constraints of the aquifers. Arsenic concentration in groundwater varies by and large between >50 and $1000 \mu\text{g L}^{-1}$, but still higher values were measured in different studies from isolated areas of the BDP. The present study suggests that arsenic concentration in groundwater is related to mobilization of arsenic from the aquifers sediments driven primarily by the reductive dissolution of the surface-reactive inorganic phases such as Fe-, Mn-, and Al-oxyhydroxides in the sediments.

Hydrogeochemical studies indicate that the groundwaters are circum-neutral (pH 6.5–7.6) with moderately to strongly reducing character (Eh 594 to -444 mV). Major ion chemistry is dominated by HCO_3 , Ca, and Mg. Dissolved organic carbon and phosphate concentrations in groundwater are high. SO_4 and NO_3 concentrations are significantly low and consistent with the high levels of dissolved Fe_{tot} and As_{tot} . HCO_3 was found to be correlated with both Fe_{tot} as well as the DOC. Aqueous speciation of As is dominated by the trivalent species and confirms the reducing character of the BDP groundwaters. Elevated concentration of arsenic is found in the aquifer sediments at different locations and depth varies between 5 and 294 mg kg^{-1} . We have also observed positive correlation between As_{NO_3} and Fe_{NO_3} ($p < 0.001$), whereas weak correlation was noted for As_{NO_3} and P_{NO_3} ($p < 0.05$). The sediments contain amorphous Fe-, Mn-, and Al-oxyhydroxides revealed by the oxalate extractable Fe ($\text{Fe}_{\text{ox}} = 4.2\text{--}174 \text{ mg kg}^{-1}$), Mn ($\text{Mn}_{\text{ox}} = 0.1\text{--}4.5 \text{ mg kg}^{-1}$), and Al ($\text{Al}_{\text{ox}} = 0.6\text{--}43.7 \text{ mg kg}^{-1}$), which accounts for the bulk of the oxalate extractable As. The mechanisms of arsenic mobilization in BDP groundwater are governed by the microbial reduction of Fe- and Mn-oxyhydroxides under anoxic conditions. Further studies are required to understand these complex biogeochemical interactions in the reducing BDP aquifers and their role in arsenic mobilization.

Arsenic enters humans via air, water, and food. High concentration of arsenic in drinking water and the food chain has caused severe health problems in rural areas of Bangladesh, especially in southern Bangladesh. The toxic character of arsenic has been known for centuries. A dose–response relationship between inorganic arsenic exposure from pumped groundwater and skin disorders (keratosis), internal cancer, respiratory, and cardiovascular problems has been established not only in Bangladesh but also for many other countries. Clinical manifestations of arsenic toxicity have also created severe social problems.

Keeping in view the poor infrastructure of the country, it is necessary to seek techniques of low-cost arsenic removal from groundwater suitable for drinking purposes in rural areas. Natural geological material such as laterite has shown promising results with an arsenic removal efficiency of 70–90%. In some areas, local materials such as coarse sand, wood charcoal, and iron filings have also been used for the removal of arsenic from drinking water. However, this is the largest single groundwater arsenic problem occurring so far in modern civilization, and it deserves special attention as well as international programs on drinking water issues.

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