Abstract

Geophysicist Glenn R. Morton has argued that a dangerous concentration of the toxic metal mercury would have been released by the Genesis Flood. This belief is based on the assumptions that the pre-Flood crust contained a concentration of mercury similar to that of the modern crust, that 90 percent of this mercury would be released as dissolved mercury ions by the Flood, and that the resulting mercury concentration of 100 parts per billion (ppb) would have been devastating to life on earth. However, a careful study of the relevant literature shows that none of these assumptions are warranted. There is good reason to believe that the pre-Flood crust contained less mercury than Morton estimated, because many mercury deposits were most likely formed by hydrothermal activity during or after the Flood. Furthermore, it is unlikely that the Flood would have fully dissolved what mercury was present, because most of it would have been in the form of highly insoluble mercury sulfide. Of the mercury that was dissolved, only 2–20 percent would be expected to be methyl mercury, which is by far the most dangerous form of the metal. It is primarily methyl mercury that accumulates in food chains and poses the greatest threat to higher life forms. There are also numerous instances documented in the literature of plant life surviving in waters with a mercury concentration higher than 100 ppb. Therefore, Morton’s challenge to the Genesis Flood does not stand; there is no reason to question the Genesis account based on mercury chemistry.

Introduction

The account of the great Flood found in Genesis 6–8 is one of the most fascinating passages in the Scriptures. In one mighty act of righteous judgment, God destroyed the world that was and created the world we see today. Many important lessons concerning God’s character and our lives can be drawn from the Genesis account of this event.

The Genesis Flood, however, is more than just a moral lesson; it was also a real historical event — one of the greatest in earth’s geological history. Today, however, many geologists dispute that such an episode could have occurred, and their concerns should be answered directly and forthrightly. If the Bible is God’s Word and therefore true, and if truth about nature can also be found via the proper application of the scientific method, then when the findings of science appear to be at odds with Scripture, there are only two possible reasons. Either my understanding of Scripture must be in error or there is a problem with the scientific evidence (Brand 1997).

With that in mind, let us examine one challenge to the veracity of the Genesis Flood — that the Flood would have released sufficient mercury to have made the survival of Noah and his family unlikely (Morton 1998). If true, it poses a serious challenge to a literal interpretation of Genesis. Conversely, if Genesis is true, that argument cannot
be. Since numerous Bible scholars have demonstrated that Genesis is presented as an historical account (Morris, 1985), then it is likely that there is a problem with the scientific basis for this argument. To evaluate that fairly, let’s start by defining the challenge.

**The Challenge**

This argument was published by geophysicist Glenn R. Morton (1998). He claims that dangerous concentrations of toxic heavy metals, particularly mercury, would have been released during the Flood. However, Genesis infers that some aquatic animals and plant life survived the Flood itself, and that the post-Flood world was habitable by Noah, his family, and the animals aboard the ark, contradicting Morton’s claim of a toxic post-Flood world. If you wish to read Morton’s argument in his own words, please see appendix 1.

Essentially, Morton argues that all of the current sedimentary rock was formed from igneous rock eroded by the Flood. During this process of erosion, he believes that 90 percent of the mercury contained in these rocks would be released. Assuming that the mercury concentration in these igneous rocks was the same as that of the modern crust, which Morton estimates at 0.1 parts per million (ppm), and that the volume of water in the Flood was essentially the same as the volume of water in modern oceans, this gives a concentration of 100 micrograms of mercury per liter of water or 100 parts per billion (ppb) for the Flood water. Morton goes on to argue that this concentration would pose a grave threat to the reestablishment of life on earth.

The basic questions we must answer are these: (1) would a global Flood really release enough mercury to create an average concentration of 100 ppb throughout the world’s oceans, and (2) would such a concentration threaten the continued existence of life afterward? To answer these questions, we must first review the basic properties and behavior of mercury in the environment.

**Mercury Chemistry**

Mercury (Hg) is a very interesting and unique chemical element. It is a liquid and releases an appreciable amount of vapor at room temperature, which is highly unusual for a metal. It will also dissolve many other metals, forming amalgams (solutions of one metal dissolved in another metal) (Greenwood and Earnshaw 1984). Due to this ability, mercury has been used to remove precious metals, such as gold and silver, from ore since as early as perhaps 2700 B.C. (Malm 1998). In fact, much of the mercury pollution that exists today in the Western United States (Pirrone et al. 1998) or South America (Malm 1998) stems from gold mining by amalgamation. Mercury amalgams have also been traditionally used as tooth fillings, although this practice has become somewhat controversial in recent years (Aposhian et al. 1995; Bjorkman et al. 1997; Jones 2004; Lorscheider et al. 1995).

Mercury pollution is a concern due to the metal’s high toxicity. This toxicity partly stems from one of mercury’s most significant properties, its ability to form especially strong chemical bonds with sulfur. Mercury tends to bind to sulfur whenever possible. Since a number of key chemical compounds within the body contain sulfur, mercury can bind to these sulfurs, preventing the compounds from performing their intended function (Silbergeld and Devine 2000; Winker et al. 2002). Within the human body, mercury can also create reactive oxygen species (Gasso et al. 2001), affect the immune system (Silva et al. 2005; Tchounwou et al. 2003; Yokoo et al. 2003), and cause birth defects (Domingo 1995; Tchounwou et al. 2003). In adults, symptoms of mercury poisoning include damage to the central nervous system (manifested as diminished fine motor skills, tremors, and memory loss), stomach pains, pain or loss of feeling in the hands and feet, vision and hearing loss, and eventually death (Auger et al. 2005; Clarkson 1998; Harada et al. 2001; Stern 2005; Tchounwou et al. 2003; Yokoo et al. 2003). In fact, the mental effects of mercury exposure were recognized long ago, although their cause was not fully understood; the phrase “mad as a hatter” refers to the tendency of hatmakers to go insane due to handling mercury, which they used to remove lint from their hats.

Mercury appears in a number of different chemical forms, including the metallic liquid Hg^0 (this is the form many readers are most likely familiar with, since it is found in older thermometers), the water-soluble ionic (an ion is a charged atom) species Hg^{2+}, the rare Hg^{2+} ion (technically an ion in which each atom has a positive one charge), and the carbon-containing molecule methyl mercury (technically CH_{3}Hg^{+} or (CH_{3})_{2}Hg, although for simplicity’s sake in this paper I will refer to all organic mercury compounds as methyl mercury). While all of these forms are toxic, methyl mercury is by far the most toxic; only a few drops on the skin can kill an adult human (Holden 1997). This is because methyl mercury in many ways behaves like an organic molecule of the type living things routinely interact with. Therefore, it is usually absorbed rather than excreted by the body (Boening 2000). Most of the major cases of mercury poisoning in recent history have involved
methyl mercury. It was methyl mercury stemming from waste dumped by a local chemical plant that contaminated fish in Minamata Bay, Japan, leading to a massive outbreak of human mercury poisoning that affected 2,262 people between 1956 and 1998, and gave mercury poisoning its clinical name of Minamata disease (Futatsuka et al. 2000). Methyl mercury, used as a fungicide to treat seed wheat sent to Iraq to stave off famine in 1971, was also responsible for the hospitalization of 6,530 people and death of 459 when hungry farmers turned the wheat into bread rather than planting it (Bakir et al. 1973). When mercury toxicity is discussed, methyl mercury is the form of most interest.

Clearly, it is not enough to simply say mercury is highly toxic, as does Morton. No matter how toxic, if mercury is not available to living things in the environment, then it does not pose a real threat. The real issue is whether a global Flood would actually release enough mercury in a sufficiently dangerous form (i.e., methyl mercury) to poison the post-Flood world. To be able to intelligently answer that question, we need to know how mercury actually behaves in natural waters.

Mercury’s behavior in an aquatic environment differs greatly between oxygen-rich (oxic) and oxygen-poor (anoxic) waters. Because the chemical reactions are different in these two environments, different chemical compounds are generated that mercury will react with. The amount of oxygen in water is, in turn, often dependent upon depth (figure 1). In general, there is more oxygen near the surface than in deeper water. Regardless of depth, mercury atoms are rarely found alone. Rather, they tend to be chemically bonded to other elements, creating a variety of compounds that will undergo different reactions.

In oxic surface waters, mercury tends to bind to the hydroxide ion (OH\(^-\)) or the chloride ion (Cl\(^-\)), although it will bind to sulfur if a sufficient amount is present (Morel et al. 1998). If a great deal of organic material (decaying plants and animals) is present and a significant amount of that material contains sulfur atoms, then as much as 95 percent of the mercury may be attached to that (Morel et al. 1998; Ravichandran 2004). In these waters, mercury is usually in the Hg\(^{2+}\) form, although it can be converted to Hg\(^0\) by bacteria or sunlight. Some bacteria possess, as a defense against mercury poisoning, a group of genes called the mer operon, which will convert Hg\(^{2+}\) and/or methyl
Once it has become Hg\textsuperscript{0}, the mercury is more toxic if the mercury concentration is too low to activate the enzyme. In general, the reaction with light is most important if the mercury concentration is low enough to activate the \textit{mer} operon; otherwise, the bacterial route dominates. The primary compound formed by the reaction of mercury with sulfide is mercury sulfide or cinnabar (Hg\textsubscript{2}S\textsubscript{2}). Most of the mercury in sediments is, in fact, found as cinnabar (Greenwood and Earnshaw 1984). Cinnabar itself is highly insoluble, so formation of cinnabar would have the net effect of removing mercury from water. Cinnabar can form more soluble compounds in the presence of other sulfides (Morel et al. 1998), elemental sulfur (Morel et al. 1998), acidic waters containing Cl\textsuperscript{−} (Mikac et al. 2003), or excess organic material (Ravichandran 2004).

Not only are sulfate-reducing bacteria partly to blame for increased mercury concentrations in anoxic waters, they are also the source of deadly methyl mercury (Morel et al. 1998). An enzyme in these bacteria will convert Hg\textsuperscript{2+} to methyl mercury (Choi et al. 1994). The methyl mercury is then absorbed by one-celled organisms, which are in turn eaten by larger organisms. In this way, the mercury moves up the food chain. For example, if 10 percent of the total mercury in an aquatic ecosystem is methyl mercury, approximately 15 percent of the mercury in tiny phytoplankton would normally be methyl mercury, increasing to 30 percent in the zooplankton that feed on the phytoplankton, and 95 percent in the fish near the top of the food chain (Morel et al. 1998) (figure 1). This can result in significant amounts of methyl mercury accumulating in the top predators, which can be toxic if eaten by humans.

Therefore, Morton greatly oversimplifies the problem. Fish have a very low uptake rate for Hg\textsuperscript{2+} and will not accumulate significant amounts of it (Morel et al. 1998). High levels of Hg\textsuperscript{2+} are of concern primarily because of its ability to be converted to methyl mercury. In fact, the U.S. Environmental Protection Agency (EPA) is considering setting new groundwater mercury standards based on the concentration of methyl mercury in the water, but on the concentration of methyl mercury in fish living in a body of water (Southworth et al. 2004). Even in the Everglades, an almost ideal environment for methyl mercury production, only 20 percent of the total mercury is methyl mercury (Cai et al. 1999). By comparison, only 3 percent of the total mercury dissolved in rivers and 2 percent in coastal ocean waters is methyl mercury (Southworth et al. 2004). One more recent study found that between 1.2 and 17.2 percent of the total mercury in peaty stream banks was methyl mercury, while less than 1 percent was methyl mercury in the soil 65 feet (20 m) from the stream (Skyllberg et al. 2003), while another found that 0.3 percent and 8 percent of the total mercury in a number of Virginia and Tennessee streams was methyl mercury (Southworth et al. 2004).

Mercury is also mobilized by the breakdown of organic material. Since these represent a major reservoir of mercury in the environment, the breakdown of peat and swamps can be a significant source of atmospheric mercury. The excess organic material helps make swamps natural mercury traps (United States Geological Survey [USGS] 2004), since this organic material will bind the mercury that evaporates into the atmosphere, be converted back to Hg\textsuperscript{2+} there, and return to earth elsewhere in rain. It is also possible for Hg\textsuperscript{2+} to be converted to methyl mercury in oxic water, but only if the concentration of Cl\textsuperscript{−} is in the right range and the correct bacteria are present (Najera et al. 2005). Oxic surface waters are not considered a major area for methyl mercury production.

Higher concentrations of mercury are often found in anoxic waters (Mason et al. 1999; Morel et al. 1998). These include deeper waters and swamps or peat bogs, where bacterial decomposition of dead plant material uses up a great deal of oxygen and poor circulation fails to replenish it. The excess organic material helps make swamps natural mercury traps (United States Geological Survey [USGS] 2004), since this organic material will bind the mercury that evaporates into the atmosphere, be converted back to Hg\textsuperscript{2+} there, and return to earth elsewhere in rain. It is also possible for Hg\textsuperscript{2+} to be converted to methyl mercury in oxic water, but only if the concentration of Cl\textsuperscript{−} is in the right range and the correct bacteria are present (Najera et al. 2005). Oxic surface waters are not considered a major area for methyl mercury production.
today's environment, they are of great importance for our
discussion of mercury and the Flood.

Geologic mercury is usually found in one of three
types of deposits: Almaden-type deposits, silica-carbonate
deposits, and hot spring deposits (Rytuba 2002). More
than one-third of all mercury mined worldwide comes
from the world's largest cinnabar deposit near Almaden,
Spain (Jebrak et al. 2002), the type of locale for Almaden-
type deposits. These ores formed from large submarine
hydrothermal vents or submarine volcanoes. Although
there is still some debate as to exactly how and when the
Almaden deposit was created, the consensus seems to be
that the main deposits were deposited from hydrother-
mal fluids as, or shortly after, the sedimentary rock layers
containing them formed, while secondary deposits were
formed by the remobilization of that mercury, perhaps in
conjunction with the introduction of new hydrothermal
mercury during later hydrothermal events (Hernandez et
al. 1999; Hugueras et al. 1999; Jebrak et al. 2002). The
layer of rock (primarily the rock quartzite) containing the
mercury has been assigned to the Silurian age (Hernandez
et al. 1999; Rytuba 1986a). This is a geological period
that young-Earth geologists believe corresponds to a time
early in the Flood (Whitmore 2007), consistent with the
Almaden volcanic/hydrothermal activity being part of the
erupting “fountains of the deep.”

Silica-carbonate mercury deposits are found along fault
zones associated with the mineral serpentinite (Rytuba
2002, 2003), which has been transformed by an influx of
carbonate and silica from low-temperature hydrothermal
fluids (Rytuba 2002; Sherlock and Logan 1995). These
fluids also contained mercury, which was deposited as cin-
nabar along fractures in the altered serpentinite (Rytuba
1986b; Sherlock and Logan 1995). These are believed to
have formed during the Tertiary, well after the Almaden
deposit (Ash 1996; Rytuba 1986b), and are often asso-
ciated with “prehistoric” mineral springs (Sherlock and
Logan 1995).

Hot spring mercury deposits form in similar fashion,
but with hotter hydrothermal fluids (Sherlock and Logan
1995). They are not associated with serpentinite (Pan-
teleyev 1996; Rytuba 2002) and often occur in environ-
ments nearer to the surface than silica-carbonate deposits
(Rytuba 2002). Some are still forming today in places like
Sulphur Bank, California (Sherlock and Logan 1995).
Although not usually classified as one of the three major
deposit types, the world’s second largest mercury deposit
occurs at Idrija, Slovenia. It is believed to have formed
in a manner similar to hot springs and silica-carbonate
deposits with hydrothermal fluids infiltrating through
already-established sedimentary rock layers and deposit-

ing mercury. However, a significant amount of mercury
also may have been in the black shale present prior to
hydrothermal enrichment (Lavric and Spangenberg 2003).
The association between mercury and black shale will be
discussed later.

With this brief overview in mind, let’s examine the first
key question relating to Morton’s challenge: would the
Flood release sufficient mercury to create an average con-
centration of 100 ppb worldwide?

How Much Mercury Would the Flood
Release?

Morton assumes that there was little to no sedimentary
rock present before the Flood and that the current volume
of sedimentary rock corresponds to the amount of igne-
ous rock crushed and redeposited by the Flood. While this
is probably an overstatement, we can use it as a maximum
for Flood erosion. Morton then assumes that Earth’s crust
contains 0.1 ppm mercury and that erosion would release
90 percent of that mercury, leading to a 100 ppb aver-
age mercury concentration in the world’s waters. Morton
considers 0.1 ppm a “very conservative” estimate, basing
this number on a USGS report (Parker 1967). This report
actually listed three values for mercury in Earth’s crust and
two for igneous rock. In both cases, the oldest value was
0.5 ppm, while newer estimates were approximately 0.08
ppm. More recent papers estimate 0.05 ppm (Lavergren

Therefore, 0.1 ppm is not conservative; if anything, it
seems a little high. Furthermore, these estimates are for the
mercury concentration of the modern crust. If the major
mercury deposits observed today (such as Almaden) were
formed during or after the Flood, pre-Flood crustal mercury
might have been much lower, especially since it appears that
they formed by mercury migrating in hydrothermal fluids
from Earth’s interior. Perhaps a better estimate would come
from looking at the mercury concentration in the bulk
silicate earth, which includes the crust and mantle. This
might better approximate the primitive mantle prior to the
formation of a true crust (Baumgardner 2000; Kargel and
Lewis 1993; McDonough and Sun 1995). This number
would be 10 ppb, or 0.01 ppm, an order of magnitude
less than Morton’s “conservative” estimate (Baumgardner
2000; McDonough and Sun 1995). However, that is only
his first error.

Morton’s 90 percent estimate of mercury mobilization
comes from a USGS paper (Siegel and Siegel 1987)
discussing mercury being released by volcanoes in Hawaii. Volcanoes are a major source of mercury; significant spikes in total atmospheric mercury correspond to volcanic eruptions (Schuster et al. 2002). Morton refers to a section of this paper that discusses mercury lost by erosion of a cooled lava flow. However, the context is the slow release of mercury from the lava flow over a century. Morton apparently assumes that the same amount would be released by rapid weathering (over the course of perhaps days) during the Flood. However, this assumption appears flawed. To understand why, let’s take a closer look at the paper Morton is referencing, one paragraph of which is quoted by Morton. Here is the paragraph directly preceding the one he quotes:

Lava flows constitute another source of mercury. Weathering brings about the slow release of soluble or solubilized constituents as the igneous materials degrade into soil minerals. Lava samples were analyzed by digestion of 100-mesh powder with 0.1 N HCl to remove soluble and loosely bound mercury. This was followed by hot 2N HNO₃ digestion to remove any mercury complex with organic ligands, and then concentrated HF to destroy the silicate matrix (Siegel and others, 1975). Samples from flows of 1840, 1923, and 1955 were obtained with the assistance of the late Gordon Macdonald; fresh Pauahi samples were collected in 1979. The results . . . suggest a 50 percent release in about 50 years and a gradual infiltration of oxidizable, presumably humic, complexing substrates (Siegel and Siegel 1987, p. 832–833).

This paragraph is followed by a table (see table 1) listing how much mercury was removed from rock samples of different ages by each of the three methods described above. The estimate of 90 percent mercury loss over the course of 100 years comes from comparing the total amounts of mercury (from all three measurement methods) for samples of different age. However, this is not the only significant thing we can learn from this data. For all the samples, the 0.1N HCl (HCl is a strong acid and normality [N] is a unit of concentration, a 0.1N solution of HCl would have a pH of 1) removed only a small amount of the total mercury. For every sample but the oldest, the 2N HNO₃ removed far less mercury than the concentrated HF.

Morton states that sedimentary rocks in the Flood were formed from mechanically crushed and chemically weathered igneous rocks, releasing mercury in the process. The igneous rocks in the study he cites were crushed, then treated with various acids to remove the mercury for analysis. This laboratory procedure was probably more chemically destructive and therefore more likely to release mercury than anything occurring during the Flood. Furthermore, it is clear that only the most extreme laboratory conditions actually removed the majority of mercury from these samples. Digestion in 0.1N HCl is a very chemically destructive process, and 2N HNO₃ is slightly more than an order of magnitude more acidic (plus more chemically reactive in other ways) than 0.1N HCl. Yet only in the sample from 1840 was more than 21 percent of the total measured mercury removed by the HCl and HNO₃ combined. In that sample, roughly 66 percent of the total mercury was removed by both procedures. The HF was successful at removing the mercury not because it is a stronger acid than the others (it’s actually weaker), but because a specific chemical reaction between the HF and silicon in the igneous rock literally breaks up the chemical structure of the rocks (this is what the authors meant by “destroy the silicate matrix”). The authors concluded that the mercury released by HCl was only loosely attached to the rock. Mercury released by HNO₃, on the other hand, was interpreted as being attached to organic compounds that had gradually entered the lava flow; HNO₃ will destroy organic material through an oxidation reaction. However, organic material presumably was not destroyed by the Flood water, nor would the Flood water have resembled

<table>
<thead>
<tr>
<th>Year lava flow formed</th>
<th>Hg extracted by HCl (ppb)</th>
<th>% of total Hg extracted by HCl</th>
<th>Hg extracted by HNO₃ (ppb)</th>
<th>% of total Hg extracted by HNO₃</th>
<th>Hg extracted by HF (ppb)</th>
<th>% of total Hg extracted by HF</th>
</tr>
</thead>
<tbody>
<tr>
<td>1840</td>
<td>5</td>
<td>3.82%</td>
<td>81</td>
<td>61.8%</td>
<td>45</td>
<td>34.4%</td>
</tr>
<tr>
<td>1923</td>
<td>16</td>
<td>4.08%</td>
<td>66</td>
<td>16.8%</td>
<td>310</td>
<td>79.1%</td>
</tr>
<tr>
<td>1955</td>
<td>10</td>
<td>0.962%</td>
<td>59</td>
<td>5.68%</td>
<td>970</td>
<td>93.4%</td>
</tr>
<tr>
<td>1979</td>
<td>0</td>
<td>0.00%</td>
<td>15</td>
<td>1.15%</td>
<td>1,290</td>
<td>98.8%</td>
</tr>
</tbody>
</table>

Table 1. The extraction of mercury from cooled lava flows (Siegel and Siegel 1987)
concentrated HF. Therefore, the digestion in HCl is probably the only process analogous to those of the Flood, and the Flood water would have been less acidic than the HCl. For the two oldest lava samples, approximately 4 percent of the mercury was removed by the HCl; in other words, only 4 percent of the mercury was readily soluble in very acidic water even after the rocks were physically pulverized, and not the 90 percent suggested by Morton!

This would not be surprising to any chemist; most of the mercury would be in the form of mercury sulfide, which is extremely insoluble. Mercury does not become a pollution problem because large amounts of mercury sulfide will dissolve in water; rather, the mercury either enters the water in a more soluble form or very tiny amounts of mercury sulfide dissolve and are then (over time) magnified up the food chain. Many studies have documented the insolubility of HgS, even in strong acids (Fernandez-Martinez and Rucandio 2003, 2005; Martin-Doimeadios et al. 2000; Mikac et al. 2003). Excess Cl− is required to make mercury sulfide soluble. How much Cl− would have been available during the Flood? At present, it is estimated that the crust only contains 185 ppm chlorine (Parker 1967), while that in the silicate earth is estimated at 17 ppm (McDonough and Sun 1995). Even the higher figure (assuming all of it converted to soluble Cl− and using the same figures Morton used to estimate the concentration of mercury in Flood water) only corresponds to 0.00586 N Cl−. This is approximately 17 times less Cl− than is present in 0.1N HCl, which was ineffective in extracting mercury. Mikac et al. (2003) noted that in the presence of 0.01N Cl−, greater than 1N HNO3 was required to extract significant amounts of mercury. The Flood water simply would not have been that acidic.

Although this fact is not mentioned by Morton, it has also been reported in the literature (Revis et al. 1989; Han et al. 2008) that a saturated solution of sodium sulfide will extract significant amounts of mercury from HgS-contaminated soils. The reality is that moderate or low concentrations of sulfide will remove mercury from water as HgS and prevent its redissolving (Piao and Bishop, 2006), while high concentrations will render it soluble again. However, using calculations such as those used for chlorine in the preceding paragraph, it is clear that even if all the sulfur in the earth’s crust was dissolved and converted to sulfide, the resulting Flood waters would contain a much lower concentration of sulfur than a saturated Na2S solution would; sodium sulfide is very soluble! Therefore, this would not significantly help Morton to get the HgS in the earth’s crust dissolved. In fact, the sulfide in the Flood waters would most likely have been at a low enough level (once the interaction of sulfur with other elements in the water is considered) to significantly decrease the solubility of the mercury. Han et al. (2008) also showed that mercury can be somewhat more easily extracted from HgS residing in soil that has been planted with crops for several seasons. However, only a small fraction of the crust destroyed by the Flood would have been topsoil involved in agriculture and much stronger acids than we would expect in the Flood waters were required to extract the mercury in that study. These findings do not make Morton’s thesis any more believable.

For the last word on this subject, Professor Donard, an environmental chemist from France who studied the extractability of mercury sulfide from the mines at Almaden, Spain, noted (Martin-Doimeadios et al. 2000, p. 365):

The total extraction results and the sequential extraction procedure have shown that mercury in the Almaden’s sediments is quite stable and presents low chemical availability. This lack of availability renders inorganic mercury methylation difficult. The results are consistent with mineralogy of mercury deposits, since cinnabar has an extremely low solubility in water, is resistant to physical and chemical weathering, and is hardly leached under acid drainage.

Although Morton’s basic thesis is clearly in error, let’s examine other ways that mercury could get into the Flood environment. First, the 40 days and nights of rain would have moved essentially all of the mercury from the atmosphere. However, this would have been insignificant: even after more than a century of industrial activity, which has dramatically increased the amount of mercury in the atmosphere (Schuster et al. 2002), there are only 6,000–10,000 tons of mercury there today (Lin and Pehkonen 1999). In the pre-Flood world, atmospheric mercury would have presumably been much less. Second, volcanoes would have released mercury, and we assume a tremendous amount of volcanic activity during the Flood. Likewise, undersea hydrothermal vents would have released mercury (Ruelas-Inzunza et al. 2003). If the “fountains of the deep” mentioned in Genesis 7:11 indicate hydrothermal activity, these might have been a significant source of mercury pollution (although for reasons I will explain later, I somewhat doubt it). Overall, these processes probably elevated the amount of mercury in the Flood environment. Still, given Morton’s
miscalculation of both the total available mercury and the percentage released into the environment, the total concentration would have been considerably less than Morton’s 100 ppb.

So what is a more reasonable estimate for the amount of mercury released by the Flood? From the outset, I want to acknowledge that this is not a simple question; as has already been shown, there would have been a number of factors affecting mercury concentration (many of which simply cannot be determined millennia later), and therefore the best that can be provided is a rough estimate. However, I believe I can at least give an estimate that is closer to reality than Morton’s. To start with, in place of Morton’s too high estimate of 0.1 ppm, let’s use Lavergren’s 0.05 ppm value (Lavergren 2005; United Nations Environment Programme 2003), which immediately cuts Morton’s number in half and equates to 50 ppb mercury in the Flood waters. Also, since Morton’s assumption that 90 percent of the mercury in the crust would be dissolved is far too high, let’s reduce that to a more reasonable value of 5 percent. (In light of the extreme insolubility of mercury sulfide, I think this is a generous estimate.) That would reduce the concentration to 2.78 ppb in the Flood waters. If we started with the bulk silicate earth mercury value of 0.01 ppm for the crust’s concentration (Baumgardner 2000; McDonough and Sun 1995), that would reduce the final value by a factor of five, giving a final concentration of 0.56 ppb mercury. Admittedly, my assumption that 5 percent of the mercury would dissolve, while in line with the published data, might be downplaying the unique properties of the Flood. If we assume that my estimate of mercury solubility is too low and arbitrarily triple it to 15 percent, that still results in mercury concentrations of less than 10 ppb, a tenth of Morton’s estimate. So I would conclude that there was between 0.5 and 10 ppb mercury in the Flood waters, with the real value probably lying closer to the smaller number. Furthermore, no more than 20 percent, and probably closer to 2 percent, of this would be the deadly methyl mercury. As we will see in the next section, while such a concentration is far from desirable, it is not catastrophic.

So Morton’s estimate of mercury released by the Flood is massively too high. However, for the sake of argument, let us assume that Morton is right and the Flood water contained 100 ppb mercury for our evaluation of the second big question: the threat such a release would actually pose to the environment.

Was Mercury a Threat to the Post-Flood World?

It is not enough for sufficient mercury to be released to yield a 100-ppb concentration in the Flood water. The mercury must have remained dissolved for enough time to have worked its way into the food chain. As noted above, the Flood water would represent a very complex chemical system. A mercury concentration near 100 ppb would have presumably activated the bacterial mer operon, so bacteria would have been converting Hg\(^{2+}\) to Hg\(^0\), which would evaporate into the atmosphere. Morton argues in his paper that Hg\(^0\) evaporation would not be an issue because the continual rainfall during the Flood would have removed it from the atmosphere. Although that would be true for the 40 days and 40 nights of rainfall, it was during this period that the Flood water was still rising, and presumably the mercury had not then reached its maximum concentration. Once the rain ended, mercury could begin to accumulate in the atmosphere, lowering the concentration in the water. However, the interplay between evaporation and precipitation would probably prevent this from being a major form of mercury removal.

There would have been a great deal of organic matter (decomposing plant and animal life) in the Flood water. This organic matter would have trapped a major amount (current studies suggest as much as 95 percent) of the mercury. If this organic bound mercury was in an oxic environment, it is unlikely that it would have been converted to methyl mercury. Thus it would pose a smaller threat because aquatic animals would not have retained Hg\(^{2+}\) to the same degree as methyl mercury. Morton does not make this distinction, simply noting that the U.S. EPA’s permissible concentration for mercury in drinking water is 2 ppb and assuming anything above this concentration would be problematic. Obviously, any significant concentration of mercury in drinking water is not good; however, it is naïve to simply assume that the EPA limit represents the absolute maximum mercury concentration above which great harm occurs. The truth is that many humans routinely drink a liquid with greater than 2 ppb mercury, namely our saliva. The amalgams used for dental fillings contain a significant concentration of mercury, and that mercury tends to find its way into saliva. One study found that people with amalgam fillings have saliva mercury concentrations anywhere from 0 ppb to 500 ppb, with an average of ~3.5 ppb before chewing and ~31.5 ppb after chewing (Ganss et al. 2000). These values were reported to be in good accord with previously published data.
values. A slightly more recent study of Hg\textsuperscript{2+} leaching from amalgams into simulated saliva reported a concentration of 15 ppb after 6 hours of contact with the amalgam and 101 ppb after 90 hours contact (Sanna et al. 2002). While it is outside the scope of this paper to discuss the controversy over the health effects of mercury dental amalgams, it is worth noting that human life is not being imperiled by ingesting concentrations of Hg\textsuperscript{2+} greater than the EPA drinking water limit and approaching Morton’s too-high estimate for the Flood water. The primary health threat from mercury does not come from ingesting water containing Hg\textsuperscript{2+}.

In an anoxic environment, on the other hand, a significant amount of methyl mercury could begin to form. This would have been the real threat: the formation and availability of sufficient methyl mercury to infiltrate the food chain and poison life. However, only a relatively small amount of the total mercury in a system is normally methyl mercury. Under ideal anoxic conditions, it would reach no more than 20 percent of the total; in less ideal conditions, it would have been only 2–3 percent of the total. So, even if the total concentration of mercury reached 100 ppb, we would only expect 2–20 ppb to exist in that dangerous form. In reality, it was probably less than that.

In a previous section, I mentioned that hydrothermal vents could have been a source of mercury contamination for the Flood. However, it is also possible they removed mercury from the system. Not only do these vents release mercury, they also release hydrogen sulfide, and sulfides react with mercury to form insoluble mercury sulfide. Therefore, excess sulfide would have removed mercury the vent released, forming cinnabar deposits. Mercury in the Flood water near the vent would also have been captured.

The Almaden mercury belt described earlier could be an example of this process. The source of mercury at Almaden was hydrothermal fluids and magma released during elevated hydrothermal activity, probably during the Flood. The mercury is believed to have been deposited at the same time that the surrounding sedimentary rock was forming or after it had formed. Since sedimentary rocks would have formed rapidly during the Flood, and since the vents were at the interface of the Flood water and the seafloor (including former continental surfaces), the vents and associated mercury were buried by Flood sediment. This is not only consistent with the geology of the Almaden deposit, but would have resulted in limiting the deposit’s contribution to contamination of the Flood water. Modern hydrothermal systems tend to be sources of mercury pollution, but during the Flood they were more likely sinks, chemically binding free mercury that was then rapidly covered by thick sediments.

Since deepwater vents would also coincide with anoxic environments, this entrapment of mercury would preferentially lock up the more dangerous methyl mercury, in my opinion. Mercury methylation in the Flood would have been occurring in deeper waters, where organic material would have been sinking down and decaying, helping to deplete the water of oxygen. With most of the mercury being bound to organic material or occurring as mercury sulfide, it would have been settling to the bottom anoxic zone as well. Normally this would generate excess mercury methylation, but in the Flood, it would have only trapped mercury in the new sedimentary rocks.

There is still debate as to the average concentration of mercury in sedimentary rock (Parker 1967; Yudovich and Ketris 2005). However, black shales are enriched with mercury, with concentrations perhaps as much as an order of magnitude above the average of other sedimentary rock (Lavergren 2005; Parker 1967; Yudovich and Ketris 2005). Also, there is still a significant debate in the geological literature as to how black shales form (Kenig et al. 2004; Lavergren 2005; Lyons and Kashgarian 2005; Schultz 2004). The consensus seems to be that they represent at least intermittently anoxic environments rich in organic material and sulfides. Such an environment would trap mercury, both in its organic material and by hydrogen sulfide generated by the sulfate-reducing bacteria expected to flourish there during anoxic periods, and promote its methylation. We know these sediments were converted into rock. And we know that these rocks contain a great deal of mercury, probably averaging between 0.4 and 0.22 ppm (Lavergren 2005; Parker 1967; Yudovich and Ketris 2005). While all the black shale seen today likely did not form during the Flood (some of it appears rather late in the geologic column for that), it is reasonable to assume that much of it did. The mercury in these shales represents a tremendous quantity that was effectively removed from the system by sedimentation. Furthermore, since the environment that most likely formed the black shales would have been conducive to mercury methylation, this would have removed the mercury most likely to be transformed into methyl mercury. Similarly, coal is often enriched in mercury (Yudovich and Ketris 2005). Rapid coal formation during the Flood would likewise bind great amounts of mercury from the environment. Please recall that the process of converting Hg\textsuperscript{2+} to methyl mercury and then infiltrating it into the
food chain in hazardous quantities is not fast. The rapid sedimentation of the Flood would have prevented this from occurring to any great extent.

A recent study (Orihel et al. 2008) supports this view. The authors added mercury spiked with various radioactive elements (so they could track the specific mercury they had added) to an experimental lake once a year for two years. They found that most of the mercury they added the first year had found its way into the sediment by the second year. A similar study (Tessier et al. 2007) using aquariums as simulated bodies of water found that most (87.9–96.2 percent) of the mercury added had found its way into the sediment within a month. Orihel and coworkers also found that most of the mercury was not converted to methyl mercury, and much of the methyl mercury that was produced remained in the sediment and was not available to be consumed by many of the organisms. This was occurring in an environment with much slower sedimentation than the flooded earth. The best evidence suggests that natural processes would have buried much of the mercury released by the Flood before it entered the food chain (figure 2).

Even in areas where mercury methylation could occur without being removed by sedimentation, there is reason to believe that net methylation of mercury would have been less than predicted. This is because the Flood would have been an “equal opportunity” polluter; the waters would have contained more than just mercury. Using Morton’s methods, we see that the same rocks contain approximately 5 percent iron (Parker 1967). The silicate earth concentration is estimated at between 18 and 19 percent (McDonough and Sun 1995). In either case, there would have been a great deal of iron in the Flood water. Studies have shown that high concentrations of iron will decrease the rate of mercury methylation (Mehrotra and Sedlak 2005). In fact, those authors suggest adding iron to wetland sediments as a way to decrease methyl mercury pollution. Between the rapid sedimentation and the high concentration of iron in the Flood water, it seems unlikely that methyl mercury formation posed a great problem during the Flood.

Morton suggests that the high concentration of $Hg^{2+}$ in the Flood water would have harmed plant life, even if it was not methyl mercury. Of course, we have already established that total mercury in the Flood water was much less than Morton’s 100 ppb, but for the sake of argument, let’s grant him that figure. To support his contention that this would have been devastating to
plant life, Morton references some data suggesting that “typically, plant-cell damage takes place with aqueous solutions containing as little as 10 [micro]g/L of Hg ion” (Morton 1998). This comes from the same paper (Siegel and Siegel 1987) on mercury and volcanoes that Morton used to get his estimate of 90 percent mercury loss during weathering. We have already seen that his presentation of that data required further scrutiny. Thus, it comes as no surprise that this one does, too.

Siegel and Siegel (1987) only mentioned this figure in passing to set up a contrast with the relatively low toxicity of Hg⁰ to plant life. They cite a paper (Siegel 1977) as the original source of the data. If we look at this original paper, we see that Morton’s figure for plant damage, while presumably made in good faith, is misleading. That is because the ten micrograms per liter value in the original paper was not referring to “plants growing in mercury-rich ground water [sic]” as Morton states, but to tobacco protoplasts. A protoplast is a cell without its protective cell wall. In this case, the author wanted to see how toxic some metals were to plant cells if they got through all the plant’s defenses and actually reached the cell. Although this is an important topic for study, it is not a very realistic analogy of how the whole organism would respond to mercury, nor was it intended by the author to be taken in that way. Still, even accepting that this study is showing the tobacco cells at their most vulnerable state, it should be noted that, while cell damage may have occurred at a 10 ppb concentration, 100.3 ppb was required to kill 50 percent of the protoplasts. In the Flood, plants did not exist as lone protoplasts, but rather as seeds or shoots that later grew into mature organisms. Therefore, a far more realistic appraisal of mercury toxicity to plants would be the concentration required to inhibit seed germination. The seed, of course, contains multiple cells with cell walls and, therefore, as Siegel stressed in this paper, is more resistant to poisoning. In fact, he reported that a concentration of approximately 802 ppb was required to inhibit the germination of 10 percent of the tobacco seeds, and that slightly over 2,200 ppb were necessary to prevent 50 percent from germinating.

This is not an isolated example. To the contrary, as Morton should have realized, the ability of plants to survive in waters containing very high mercury concentrations has been reported many times in the literature. For example, a more recent paper concerning mercury toxicity to tobacco states, “Tobacco has been shown to be highly resistant to environmental mercury, accumulating up to 5,000 µg/g under chronic low-level exposure with no symptoms of toxicity” (Suszcynsky and Shann 1995). This means the tobacco can accumulate within it 5,000 ppm mercury without seeing any ill effect. Now, one might argue that this is only referring to chronic low-level exposure, not a sudden high-level exposure such as the Flood could cause. The paper addresses this as well. The author submerged the roots of tobacco plants in solutions with varying concentrations of mercury and observed the effects on the plants. Keep in mind that Morton’s estimate for mercury in the Flood water (far too high to begin with) is only 100 ppb or 0.1 µg/ml. Suszcynsky and Shann (1995, p. 65) noted:

An inhibition of whole plant growth was demonstrated in plants whose roots were exposed to HgCl₂ with the severity of inhibition corresponding to increasing treatment levels. Plants at higher treatment levels (>10.0 µg Hg²⁺/ml) displayed minor visible symptoms of toxicity (chlorosis) but no tissue death. . . . Although growth in plants exposed to lower treatment levels (≤ 1.0 µg Hg²⁺/ml) was slightly inhibited, there was no apparent threat to their viability as that which occurred in plants exposed to higher treatment levels.

In other words, the survival of the tobacco plants was not threatened by a mercury concentration ten times higher than Morton’s estimate, and there was no actual tissue death in plants to a concentration ten times higher than that! Tobacco is not the only plant resistant to mercury. Another study reported that 47 percent of alfalfa seedlings exposed to an approximately 600 ppb Hg²⁺ solution for 24 hours showed no ill effect, and 11 percent showed no effect after 24 hours’ exposure to 6,000 ppb mercury (Ortega-Villasante et al. 2005). Seedlings from two types of rice were shown to undergo slightly over 50 percent germination in a solution of more than 20,000 ppb mercury, while the germination rate was between 70 and 80 percent in a solution a tenth as concentrated (Mishra and Choudhuri 1999). While these may have been far from healthy rice plants, they were surviving in waters that contained orders of magnitude more mercury than even Morton suggests the Flood would have contained. A study of the aquatic plant Vallisneria Spiralis did not note ill effects on plants from exposure to any concentration less than approximately 200 ppb mercury (Gupta and Chandra 1998). Water lettuce is reported as surviving, albeit with complete inhibition of new root formation, for three weeks (the plants did not all die at the end of three weeks, the study just ended) in solutions of
Appendix 1: Morton in His Own Words

R. Morton [this is me] “Prolegomen to the Study of the Sediments,” CRSQ, Dec. 1980, p. 162–167). All of this material must have come from igneous rock.

Given that igneous rocks are around 3.3 g/cc (3300 kg/m^3) and sedimentary rocks are around 2.5 g/cc, we can correct for this and we find a .75 reduction factor to put the sedimentary rocks back to igneous. Thus, 477 x 10^6 km^3 or 4.77 x 10^17 cubic meters of igneous rocks must have been eroded.

An earlier version of this note was criticized for not making explicit an assumption. The assumption is this. Within the YEC model, the prediluvial rock, which mostly would have been granite and basalt, must have been mechanically crushed, and then rapidly altered chemically to separate the feldspar and quartz fractions. Only in this way can the vast quantities of sand and shale seen in the sedimentary rocks be explained. For God to have created the vast quantities of sand and shale on the primeval earth would be a case of God deceptively creating the appearance of age when no such appearance would be needed. On the primeval earth, only a thin layer of soil would be required, not 40 to 60,000 feet of it. By the process of mechanical crushing and rapid chemical weathering, much of the mercury contained in the rock would have been released. This is consistent with what is known to occur in the weathering of basalts in which 90 percent of the mercury in the basalts is released to the environment in about a century.

“If Kilauea lava typically cools with about 1,000 [micro] g/kg of mercury and proceeds to release 90 percent, then this still constitutes only a minor source of the element. The 1840 eruption produced about 400 x 10^6 m^3 of lava weighing perhaps 16 x 10^9 kg. Thus this lava contained a total of 16 x 10^6 g (16 tons) of mercury, of which about 14 tons was released in about a century. In contrast, Halemaumau yields 260 tons annually when it is not erupting” (Siegel and Siegel 1987, p. 833).

What is the mercury content of the crust of the earth? Using the very conservative value of .1 parts per million (ppm) we find that the flood would have ground up and released .0000001 * 4.77 x 10^17 cubic meters x 3300 kg/m^3 x .9= 1.4 x 10^14 kg or 1.4 x 10^17 g or 1.4 x 10^23 micrograms. I place this in all three units because of the need below.

All of this would have been released into the oceans for the fish to ingest. The volume of the ocean is 1.4 x 10^21 liters. So the amount of mercury in a liter is 1.41 x 10^23 micrograms/1.4 x 10^21 liters = 100 micrograms per liter of water.
How bad is it? Consider this: “Typically, plant-cell damage takes place with aqueous solutions containing as little as 10 [micro]g/L of Hg ion” (Siegel and Siegel 1987, p. 830).

An expert might question the relevance of this fact to the Flood since the above refers to plants growing in mercury-rich ground water [sic]. Since plants were not taken on the ark, they must have survived by floating on the surface of the flood waters. And many young-earth creationists have suggested that such vegetable mats were responsible for the coal bed formation. Thus, the damage which mercury laden waters would cause to these floating plants is something that must be addressed by global flood advocates.

The EPA does not allow more than 2.4 micrograms/liter, which is the EPA’s Critical Maximum Concentration for fresh water discharge from an industrial site. This was set up to protect aquatic life from deleterious effects from mercury (Gray and Sanzolone 1996, p. 5).

How about for animal ingestion? This was found on the Internet: “The EPA has set a limit of 2 parts of mercury per billion parts of drinking water (2 ppb = 2 micrograms/liter). The EPA requires that discharges or spills of 1 pound or more of mercury be reported” (http://atsdr1.atsdr.cdc.gov:8080/tfacts46.html).

For those who don’t know, 100 micrograms per liter is 50 times more than the EPA would allow for an anthropogenic release. I guess the EPA would initiate regulatory action against Noah’s Flood for polluting the oceans.

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