ASSESSING THE RESPONSE OF EMERALD LAKE, AN ALPINE WATERSHED IN SEQUOIA NATIONAL PARK, CALIFORNIA, TO ACIDIFICATION DURING SNOWMELT BY USING A SIMPLE HYDROCHEMICAL MODEL

By Richard P. Hooper, Christopher T. West, and Norman E. Peters

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CONVERSION FACTORS

For those readers who may prefer to use inch-pound units rather than metric (International System) units, conversion factors for the terms used in this report are listed below:

<table>
<thead>
<tr>
<th>Multiply metric unit</th>
<th>by</th>
<th>To obtain inch-pound units</th>
</tr>
</thead>
<tbody>
<tr>
<td>millimeter (mm)</td>
<td>0.03937</td>
<td>inch (in.)</td>
</tr>
<tr>
<td>meter (m)</td>
<td>3.281</td>
<td>foot (ft)</td>
</tr>
<tr>
<td>kilometer (km)</td>
<td>0.6214</td>
<td>mile (mi)</td>
</tr>
<tr>
<td>hectare (ha)</td>
<td>2.471</td>
<td>acre</td>
</tr>
<tr>
<td>square meter (m²)</td>
<td>10.76</td>
<td>square foot (ft²)</td>
</tr>
<tr>
<td>liter (L)</td>
<td>0.2642</td>
<td>gallon (gal)</td>
</tr>
<tr>
<td>cubic meter (m³)</td>
<td>35.31</td>
<td>cubic foot (ft³)</td>
</tr>
<tr>
<td>cubic meter per day (m³/d)</td>
<td>264.2</td>
<td>gallon per day (gal/d)</td>
</tr>
<tr>
<td>liters per second (L/s)</td>
<td>0.0353</td>
<td>cubic feet per second</td>
</tr>
</tbody>
</table>

In this report, “sea level” refers to the National Geodetic Vertical Datum of 1929 (NGVD of 1929)—a geodetic datum derived from a general adjustment of the first-order level nets of both the United States and Canada, formerly called Sea Level Datum of 1929.
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ABSTRACT

A sparsely parameterized hydrochemical model has been developed by using data from Emerald Lake watershed, which is a 120-hectare alpine catchment in Sequoia National Park, California. Greater than 90 percent of the precipitation to this watershed is snow; hence, snowmelt is the dominant hydrologic event. A model which uses a single alkalinity-generating mechanism, primary mineral weathering, was able to capture the pattern of solute concentrations in surface waters during snowmelt. An empirical representation of the weathering reaction, which is based on rock weathering stoichiometry and which uses discharge as a measure of residence time, was included in the model. Results of the model indicate that current deposition levels would have to be increased between three- and eight-fold to exhaust the alkalinity of the lake during snowmelt if there is a mild acidic pulse in the stream at the beginning of snowmelt as was observed during the study period. The acidic pulse in the inflow stream at the onset of snowmelt was less pronounced than acidic pulses observed in the meltwater draining the snowpack at a point using snow lysimeters or in the laboratory. Sulfate concentrations in the streamwater were the most constant; chloride and nitrate concentrations increased slightly at the beginning of snowmelt. Additional field work is required to resolve whether the an acidic meltwater pulse occurs over a large area as well as at a point (implying sulfate-regulating mechanisms in the soil) or whether, due to physical and chemical processes within the snowpack, the acidic meltwater pulse is attenuated at the catchment scale. The modest data requirements of the model permit its applications to other alpine watersheds that are much less intensively studied than Emerald Lake watershed.

INTRODUCTION

The Emerald Lake watershed, an alpine catchment in Sequoia National Park in the Sierra Nevada of California, has been the subject of an intensive 5-year research effort under the Integrated Watershed Study (IWS) of the California Air Resources Board. All aspects of the watershed—its hydrology, geology, soils, vegetation and aquatic biota—were studied to gain an understanding of the sensitivity of the ecosystem to acidification (California Air Resources Board, 1986). Because over 90 percent of the precipitation in this watershed is in the form of snow, the emphasis of the field program was on the response of the catchment to snowmelt. Field data were collected starting in October, 1985, but data from 1986 and 1987 are more complete than from the first year and they reflect the refinement of snow-measurement techniques from information gained during the first year of sampling.
Purpose and Scope

One phase of the IWS was the development of simulation models to predict surface-water chemistry. These models were to serve as research tools to integrate information gained from the field program and to identify gaps in understanding the chemical mechanisms controlling surface-water chemistry, and as predictive tools to provide a means for forecasting the chemical response of the watershed to different depositional loadings of atmospheric acidity. This paper presents a lumped, sparsely parameterized simulation model, called the Alpine Lake Forecaster, or ALF, that was developed from the IWS data. ALF is a model of intermediate complexity when compared with a fully distributed model (Sorooshian and others, 1989) and a regional lake acidification model (Nishida and others, 1989), which also were developed from this data set.

ALF was designed and calibrated from data collected during water year 1986, which extended from October 1, 1985, to September 30, 1986. This year was an exceptionally heavy snow year, with twice the usual snow accumulation, and a maximum snow depth of more than 5 meters (m). Water-quality samples of surface water were taken on a biweekly basis from the onset of snowmelt throughout summer and fall (Dozier and others, 1989). Intensive sampling of infrequent summer rainstorms was performed only opportunistically (Melack and others, 1989). Snow samples were taken on an event basis during winter, and a whole-basin survey was performed at maximum snow accumulation to estimate the water volume and ionic loadings to the catchment. Data from 1987, a light snow year with one-quarter the amount of snow as 1986, were used for validation of the model. The sampling regime was for surface water samples was similar to that in 1986. Based upon the experience gained from the previous year, only a snow survey at maximum accumulation was performed; no event snow samples were collected. Data considered in construction of the model included stream- and lake-water quality and quantity, snowpack quality and quantity, soil column studies, and geochemical characterizations of the bedrock.

Site Description

The following site description, except where noted, is excerpted from Dracup and others (1989) to which the reader is referred for more details. The Emerald Lake watershed (36° 35' 49" N, 118° 40' 30"W) is within Sequoia National Park on the western slope of the Sierra Nevada in the Marble Fork of the Kaweah River drainage, approximately 5 kilometers (km) east-southeast of Lodgepole (fig. 1). The area of the watershed is about 120 hectares (ha) and altitudes range from 2,800 m at the lake outlet to 3,416 m at the top of Alta Peak. The watershed is an alpine cirque that has the typical bowl shape and more than 180 degrees of aspect. The long axis of the watershed (approximately 1.7 km in length) is oriented from southeast to northwest, and more than half of the watershed has a north to northeast aspect (fig. 2).

Bedrock underlying the Emerald Lake watershed consists of granite, granodiorite, and aplite (Clow, 1987). The granite consists of 2 to 7 percent biotite and hornblende by volume; however, the percentage of these more readily weathered minerals in the granodiorite ranges between 12 and 17 percent by volume. More than one-third of the basin consists of exposed bedrock, and most of the remainder is covered by a thin mantle of talus or colluvium. Only 20 percent of the basin is covered by soils (Lund and others, 1987). Bedrock comprising the basin floor and cliffs has been exposed recently by glacial erosion and frost action and, thus, is weathered only slightly. Both subalpine and alpine vegetation are present; western white pine (Pinus monticola) is the predominate tree species and willows (Salix sp.) are the dominant shrub.
Figure 1.—Map showing location of study site.
Figure 2.—Map showing subbasins within Emerald Lake watershed.
Precipitation is seasonal in the alpine regions of Sequoia National Park. Stephenson (1988) estimated that 18 percent of the annual precipitation falls from September through November, 50 percent from December through February, 30 percent from March through May, and only 2 percent from June through August. Snowmelt begins in March or April and continues through June, except in the heaviest snow years when melt continues through August. Consequently, few rain-on-snow events occur. More commonly during late spring and early summer, snowstorms occur after the melt has begun.

Emerald Lake is a tarn fed by three major and five minor stream channels; only inflows 1, 2, and 4 (fig. 2) continue to flow after the snow has melted in their respective drainage basins. Two of these perennial or nearly perennial streams (inflows 1 and 2, fig. 2) are split parts of a larger stream that drains along the main axis of the basin; this drainage accounts for 60 percent of the total inflow area. For modeling purposes, the drainage area of inflows 1 and 2 was designated as a single unit. Outflow from the watershed seems to occur only through the lake outlet. The geomorphometry of the lake suggests that the low, transverse ridge of bedrock that runs along the master joint trace on the valley floor dams the lake and is impermeable. There is no visual evidence of ground-water seepage at land surfaces downslope of the lake. Seepage losses from the watershed are assumed to be negligible.

**Background**

Alpine watersheds pose a number of challenges to the construction of hydrochemical models. Snow accounts for the vast majority of the hydrologic inputs to these systems, and therefore, snowmelt and not rain, is the important hydrologic event to be modeled. Previous models, such as the Birkenes model (Christophersen and others, 1984) and the Integrated Lake-Watershed Acidification Study (ILWAS) model (Chen and others, 1983), have used degree-day formulations to predict snowmelt. These models are appropriate for forested catchments where solar radiation and wind are not substantial components in the energy budget of the snowpack (Miller, 1950). In sparsely vegetated alpine watersheds, a more complete evaluation of the energy budget is required, but measurement of the necessary radiation values is logistically difficult. A major part of the field effort at Emerald Lake was devoted to obtaining these data, which have been used in an energy balance model of snowmelt at one point (Marks, 1988). The application of the point model to the complex terrain of the Emerald Lake watershed is an area of active research (Dozier, 1988).

Even with the prediction of meltwater generation at the surface of the snowpack, routing of water through the pack is difficult to simulate. In watersheds for which models have been developed (for example, the Adirondacks, southern Norway) and where snowpacks are on the order of 1 m deep, observations can be made as to whether the majority of snowmelt infiltrates the soil (i.e., there is no basal ice layer) or flows overland. Such observations are difficult, if not impossible, to make in alpine terrain; thus water movement through the pack is not well understood and modeling this process remains a matter of conjecture.

Moreover, existing process-oriented hydrochemical models (the ILWAS and Birkenes models) rely on a hydrologic accounting scheme that requires an accurate measure of the timing and quantity of the entry of snowmelt or rain into the watershed. Predicting or measuring snowmelt is much more difficult than rainfall, particularly in deep snowpacks. Any error in the estimates of timing and magnitude of meltwater generation will lead to errors in parameterization of the hydrologic part of the model, such as the capacity of soil layers or the rates of release from the soil layers (for example, their conductivity or other rate constant). Thus, errors in the hydrologic model can influence chemical parameterization and predictions.
A hydrologic accounting model also requires that a hydrologic budget, which divides the water input to the watershed into outflow, evapotranspiration losses and change in storage within the watershed, can be constructed with little error. For water year 1986, an acceptably precise hydrologic budget (less than 5 percent error) was obtained for the entire melt period (Dozier and others, 1989). However, when finer time steps were considered, the errors were too large for modeling purposes. A simple extrapolation of the point-snowmelt model (Marks, 1988) to the entire watershed yielded incorrectly timed meltwater inputs. Furthermore, the water yield from the snowpack, as determined from sequential snow surveys, was inconsistent with the measured lake outflow (Dozier and others, 1989). This lack of agreement probably was due both to errors in the snow survey, which were exacerbated owing to the great depth of the pack, and to errors in the stage-discharge relation, which had to be extrapolated to high stage values that were beyond the range of calibration. The difficulties in achieving an acceptable hydrologic budget led to consideration of a simpler model structure that did not depend on a hydrologic budget, but that could still be used to achieve the objectives of the modeling program.

Acknowledgments

The authors would like to thank K.A. Tonnessen of the California Air Resources Board for her persistence and organizational abilities. J.M. Melack and Jeffrey Dozier provided useful criticisms of the model and insights into the hydrologic and chemical mechanisms operating at Emerald Lake. Mark Williams, Richard Kattelmann, and J.O. Sickman responded to our many data requests and assisted in the interpretation of these data. This work was performed in cooperation with the California Air Resources Board.

MODEL STRUCTURE

ALF consists of a coupled hydrologic and chemical model. The hydrologic model, which provides the basic structure for the model, determines the quantity of discharge from each terrestrial subbasin to the lake and the volume of the lake epilimnion into which the streamwater mixes. The chemical model calculates the equilibrium concentration of the solutes for the streamwater and the lake water based upon primary mineral weathering and dissolution of carbon dioxide. These models are described in more detail in the following sections.

Hydrologic Model

To eliminate the need for a hydrologic accounting model, discharge was assumed to equal snowmelt on a daily basis. This assumption implies that there is no change in lake or ground-water storage during the snowmelt period, which is not so unrealistic as it may seem. The lake stage varies between 1 and 2 millimeters (mm) on a weekly basis (Richard Kattelmann, Univ. of California, Santa Barbara, written commun., 1987), which is equivalent to a volume of approximately 50 cubic meters (m³). Daily outflow from the lake during snowmelt ranges between 1,000 and 35,000 m³. Ground-water storage in unconsolidated surficial deposits has been estimated at 100,000 ± 50,000 m³ (Dracup and others, 1989). During the first part of snowmelt, some meltwater recharges the ground-water system, which is depleted during winter. A schematic of this simple hydrologic model is shown in figure 3. The total daily runoff from the terrestrial part of the watershed is set equal to the measured daily discharge from the lake.
Because the snowpack melts at different rates depending on the aspect and slope of the land surface, the watershed was divided into seven subbasins (fig. 2). The proportion, \( P_i(t) \), of total discharge, \( Q(t) \), for each subbasin \( i \) on day \( t \) is determined by the relative amount of solar radiation to that subbasin:

\[
P_i(t) = \frac{r_i(t)A_i}{\sum_{i=1}^{n} r_i(t)A_i}
\] (1)

The variable \( r_i \) is the potential solar radiation per unit area as determined by an algorithm of Swift (1976), and it is a function of latitude, slope inclination, aspect, and day of year. The variable \( A_i \) is the snow-covered area of the \( i \)th subbasin; and \( n \) is the total number of subbasins in the watershed. Each subbasin is credited at the start of the snowmelt season with part of the total seasonal discharge from the lake during the entire snowmelt period based on the proportion of the subbasin area; and each subbasin meets the discharge demand (i.e., \( P_i(t)Q(t) \)) as long as enough water remains in the subbasin. After the subbasin depletes its store of water, the discharge is apportioned among the remaining subbasins. If the snow-covered area is not known, the subbasin area can be used, although this tends to overestimate the contribution from subbasins with relatively less snow-covered area than other subbasins. Snow-covered area was measured at Emerald Lake, but the values by subbasin were not available in time for this modeling exercise; therefore, the subbasin area was used for the results presented in this report. The values required for the application of equation (1) to the seven subbasins are shown in table 1.
Table 1. Characteristics of terrestrial subbasins

[10^3 m^2, thousands of square meters]

<table>
<thead>
<tr>
<th>Subbasin</th>
<th>Area (10^3 m^2)</th>
<th>Percent of total area</th>
<th>Slope (degrees)</th>
<th>Aspect (degrees)</th>
</tr>
</thead>
<tbody>
<tr>
<td>West joint</td>
<td>140</td>
<td>12</td>
<td>34</td>
<td>43</td>
</tr>
<tr>
<td>SW gully</td>
<td>133</td>
<td>11</td>
<td>31</td>
<td>27</td>
</tr>
<tr>
<td>Inflow 3</td>
<td>141</td>
<td>12</td>
<td>28</td>
<td>10</td>
</tr>
<tr>
<td>Inflow 1&amp;2</td>
<td>396</td>
<td>33</td>
<td>34</td>
<td>305</td>
</tr>
<tr>
<td>NE gully</td>
<td>245</td>
<td>20</td>
<td>27</td>
<td>275</td>
</tr>
<tr>
<td>Pinestand</td>
<td>135</td>
<td>11</td>
<td>34</td>
<td>253</td>
</tr>
<tr>
<td>Meadow</td>
<td>9.28</td>
<td>0.8</td>
<td>14</td>
<td>327</td>
</tr>
</tbody>
</table>

Because there is not an accounting model with explicit reservoirs, routing of rainfall that occurs during the melt period had to be determined from the discharge. If the discharge increased on a day when rain was recorded, all of the increase in discharge was assumed to be rain, if there was a sufficient volume of rain. If not, the volume unaccounted for was supplied from the snow pack. If more rain fell than the discharge increased (as was usually the case), the “excess” rain was areally proportioned among the subbasins and added to the snowpack. Rainfall volume was minor in 1986 but provided more than 10 percent of the precipitation in 1987.

The thermocline and ice depth in the lake are determined by linearly interpolating the field values that were measured monthly in winter and biweekly during the melt period and summer. The depths are converted to volumes by using the lake bathymetry provided by J.O. Sickman (Univ. of California, Santa Barbara, written commun., 1987). The ice layer is assumed to be completely contained within the epilimnion, thus reducing the volume of this layer. The volumes of all three layers for the years that were modeled are shown in figure 4. Streams from the terrestrial part of the watershed flow into the epilimnion and the lake outflow is from the epilimnion; the hypolimnion is hydraulically separated from the surface water, and does not contribute to streamflow. Each layer is assumed to be completely mixed.

This simple hydrologic model structure eliminated artificial parameters that would have had to be fitted to the data, as is the case with the previously mentioned hydrochemical models. As long as the assumption holds that there is no change in storage of water in the lake or in the aquifers, this hydrologic model will provide a more powerful test of the proposed chemical mechanisms, in terms of a hypothesis test, than a model that contains artificial parameters.
Figure 4.—Volume of ice layer, epilimnion and hypolimnion, 1986 and 1987.
**Chemical Model**

The chemical reactions that are contained in ALF can be divided into those which generate alkalinity, those which are part of the biological nitrogen cycle, and others which include carbonate buffering and non-reactive solutes. The representation of these reactions included within ALF are described in the next three sections. The description of the chemical model concludes with a discussion of the numerical techniques used to solve to simultaneous equations which constitute the chemical model.

**Alkalinity generation**

The dominant sources of alkalinity in watersheds with young soils that contain fresh mineral faces and few sesquioxides, are primary mineral weathering and cation exchange (Drever, 1988, p. 207-231). Weintraub (1986) characterized the mineralogy and chemistry of soils, surficial material, and bedrock from Emerald Lake, and conducted laboratory leaching studies of undisturbed soil columns. Weintraub (1986) concluded that weathering was the dominant alkalinity-generating mechanism in the watershed. Lund and others (1987) further characterized the soil chemical characteristics and they concluded that the Emerald Lake soils are acidic (pH from 4.4 to 4.8) and have very low percent base saturation, which may only provide limited buffering of acidic inputs. In addition, the interrelation of solute compositions of streamwaters were evaluated by Lund and others (1987) from which they concluded that primary mineral weathering is a primary source of alkalinity. We, therefore, included mineral weathering in the model but did not include cation exchange.

The mineralogy of the bedrock, which determines the rate and stoichiometry of primary weathering, is dominated by quartz and feldspar and contains minor amounts of biotite and hornblende (Weintraub, 1986). A variety of secondary vermiculite minerals were identified in Emerald Lake soils (Weintraub, 1986). Incongruent weathering of aluminosilicate minerals produces secondary minerals, base cations, and dissolved silica (SiO₂) either while consuming hydrogen ion or releasing alkalinity. In the absence of detailed mineral-weathering information, the stoichiometry of the mineral weathering was simplified by combining the individual aluminosilicate minerals into one reactant called rock. Rather than tracking each base cation individually, the base cations (Na⁺, K⁺, Ca²⁺, and Mg²⁺) were combined as the sum of base cations, herein called SBC. The slope of the least squares regression of SBC concentration (in microequivalents per liter (µeq L⁻¹)) on dissolved silica (in micromoles per liter(µmol L⁻¹)) for the 1986 inflow data (fig. 5) is 1.2 (r² is 0.77), and therefore, the following incongruent dissolution reaction is consistent with these data

\[
\text{Rock} + 1.2\text{H}^+ \rightarrow 1.2\text{SBC}^+ + \text{SiO}_2 \text{(aq)} + \text{secondary mineral.} \quad (2)
\]

This reaction, however, is subject to kinetic constraints. The hydrologic model does not permit an explicit calculation of the residence time of water within the terrestrial system, so a standard kinetic model cannot be fit to the data. Instead, we use the lake discharge, averaged over the 10 days prior to the sample collection, as a surrogate for residence time. A regression of the log of the reaction constant, K, where
FIGURE 5.—Sum of base cations and silica concentration in streamwater samples collected from all sampled inlets to Emerald Lake in 1986.

\[ y = 2.0 + 1.2x \quad R^2 = 0.77 \]

\[ K = \frac{[\text{SBC}^+]^{1.2}[\text{SiO}_2]}{[\text{H}^+]^{1.2}} \]  
\hfill (3)

on the average discharge (fig. 6) yields the following equation,

\[ K = 0.0082 \times 10^{-0.000044Q'} \]  
\hfill (4)

where \( Q' \) is the 10-day average discharge. Equation 4 explains 80 percent of the variation in the data. Because this is the central equation in the model, two different intercepts, 0.005 and 0.02, that bracket the observations were used to assess the sensitivity of the results to this equation (fig. 6). Model results based on the lower line are referred to as the “lower bound” case because these results have lower concentrations of alkalinity, SBC, and silica than the “upper bound” case. The watershed is more sensitive to acidification under the lower bound case because less of the hydrogen ion is neutralized by the weathering reaction.
Nitrogen cycle

A simplified representation of the nitrogen cycle is included in the model. Ammonium concentration in the main inflow is very low, generally much less that 1 μeq/L. Therefore during biotically active seasons, all ammonium is assumed to be taken up by the biota; each mole of ammonium is exchanged for one mole of hydrogen ion. In addition, a part of the nitrate, \( v \), also is retained along with an equivalent amount of hydrogen ion. When the biota are not active, all the ammonium is oxidized to nitrate, which is passed through the system, and to two moles of hydrogen ion; no nitrate is taken up by the biota. After biotic uptake, if any, nitrate is treated as a conservative species. We assumed that the biota were active throughout the snowmelt period and set \( v \) to 0.10 (that is, 10 percent of the nitrate is retained by the biota).

Other chemical species

A standard representation of the carbonate buffering system is included. When the lake is covered by ice, the system is assumed to be closed to the atmosphere; during the lake’s ice-free season and during all times in the terrestrial part of the watershed, the system is assumed to be open to the atmosphere. The partial pressure of carbon dioxide (CO₂), to which water equilibrates in the open system, is a parameter in the model to be fit. Chloride and sulfate are treated as conservative species in the model.
Solution technique

A modified Newton's method (Morel and Morgan, 1972) was used to solve the system of nonlinear simultaneous equations that describe weathering and carbonate buffering in the terrestrial subbasins. There are four variables in the system, hydrogen ion, bicarbonate, silica, and SBC, that are determined from charge balance constraint, weathering equilibrium, weathering stoichiometry, and carbonate equilibrium. Temperature corrections are made for the carbonate system and for the hydrolysis of water; activity corrections are not made owing to the low ionic strength of the surface waters. The mathematical solution seems to be identical to an equilibrium solution, yet the equilibrium “constant” for the weathering reaction changes as discharge changes. These equations are solved on a daily time step by using the input concentration of the meltwater. On days when rain fell, the “excess” rain, whose chemistry is an input to the model, is assumed to react with the rock and then mix with the meltwater which has also reacted with the rock. This mixture is then re-equilibrated before flowing into the lake.

Within the lake, no chemical reactions other than carbonate buffering occur. The volume of the epilimnion is increased by inflow from the terrestrial system, thinning of the ice layer, and deepening of the thermocline. The appropriate quantities of solutes are advected with the water, and the system is re-equilibrated assuming that the epilimnion is completely mixed. The time step concludes with water of the same quality as the epilimnion flowing out of the lake.

Within the chemical model, there are only five parameters that must be calibrated: the two regression coefficients for the weathering reaction (equation 4), that part of nitrate retained by the biota (v), the date at which the biota become active, and the partial pressure of CO2. This latter parameter must be fitted because surface waters frequently contain more dissolved CO2 than would be predicted from equilibrium with atmospheric levels of CO2. Typical partial pressures range from two to five times the atmospheric partial pressure of CO2. Dissolved inorganic carbon was not measured in surface-water samples collected from Emerald Lake. The equilibrium constants and enthalpy values (used for temperature correction) for the carbonate system and for water hydrolysis are considered to be well known and are not subject to calibration.

Model Inputs

ALF was designed to utilize data collected for the Integrated Watershed Study. However, its input requirements are modest enough to allow its application to less intensively studies sites. Table 2 shows the hydrologic and chemical inputs necessary to run the model. The most critical measurements are the daily discharge from the lake and the mass of solutes contained in the snowpack at maximum accumulation. Rain water quality primarily affects stream and lake water quality on the day which the event happens. However, if there is any “excess” rain as defined above, the solutes associated with this rain are incorporated into the snowpack. Thermocline depth and ice thickness determine the hydraulic mixing volume of the lake and are of secondary importance. Water temperatures are used to correct thermodynamic equilibrium constants and also are not critical. The remaining inputs are used to determine initial conditions for the model.
Table 2. Model Inputs

<table>
<thead>
<tr>
<th>Hydrologic</th>
<th>Chemical</th>
</tr>
</thead>
<tbody>
<tr>
<td>Daily discharge from lake</td>
<td>Total mass of solutes at maximum</td>
</tr>
<tr>
<td>Thermocline depth</td>
<td>snowpack accumulation</td>
</tr>
<tr>
<td>Ice thickness</td>
<td>Rainwater quality</td>
</tr>
<tr>
<td>Temperature of inflows</td>
<td>Initial lake water quality</td>
</tr>
<tr>
<td>Temperature of epilimnion</td>
<td>Initial chemical composition of ice on lake</td>
</tr>
</tbody>
</table>

1 May be interpolated from field measurements to derive daily values necessary to run model.

Although the mass of solutes in the snowpack is an input to the model, the meltwater quality during the snowmelt must be calculated. The first meltwater from the snowpack typically contains a high concentration of solutes (Johannessen and Henriksen, 1978). This first “pulse” of solutes from acidic snowpacks is believed to be the cause for the sharp pH depressions during the spring snowmelt (Skartveit and Gjessing, 1979), and has been linked to fish kills in Norway (Leivested and Muniz, 1976). Therefore, the intensity of the acidic pulse is a key factor in determining the impact of acidic pollutants contained within the snowpack on aquatic biota. Both the Birkenes and ILWAS models assume an exponential decline in the release of solutes from the snowpack. Specifically, the concentration factor (the ratio of the concentration of the meltwater to the bulk concentration of the snowpack) is assumed to decline exponentially. Because the pattern of solute release is so critical, particularly when considering different depositional scenarios, we chose to parameterize the solute release differently. An elution rule in the form of $x$ percent of the solutes released in the first $y$ percent of the meltwater (for example, 80/20 rule means 80 percent of the solutes are released in the first 20 percent of the meltwater) and a maximum concentration factor for the first day (the ratio of concentration on that day to the average concentration of the solute in the snowpack) are specified. Two exponential curves, one for the period when the first $y$ percent of the water drains from the pack and a second curve for the remainder of the melt period, define the meltwater concentration. These curves meet the specified elution rule and concentration factor along with a continuity condition that the concentration does not change abruptly between the two curves. The results of this approach are compared with the experimental data of R. C. Bales (Univ. of Arizona, written commun., 1988) on a cumulative basis in figure 7. All solutes are treated identically; no provisions are made for “preferential” elution (that is, different solutes being released at different rates), which has been reported by Brimblecombe and others (1987).
Figure 7.—Laboratory and modeled solute release from snowpack.

Because snowpacks melt at different rates in each subbasin owing to different radiation inputs, meltwater concentration curves are calculated for each subbasin. Consequently, ionic pulses from subbasins where snowmelt occurs later in the season are contributed to surface waters after earlier ionic pulses from subbasins where snowmelt has already occurred. This distribution of meltwater generation leads to a much less intense ionic pulse to the lake than if the entire snowpack melted as one unit. The total ionic loading in the snowpack for all ions was estimated by Dozier and others (1987) from field observations (table 3). An artificial solute called was created to create an exact charge balance. Typically there is a small cation charge excess which arises from the unmeasured contribution of organic anions to the charge balance of the solution. If this solute were not created, the charge balance constraint in the chemical model would adjust the hydrogen ion concentration to meet the charge balance. The concentration of this artificial solute was always very small.
MODEL RESULTS

Errors in the prediction of one charged solute will affect predictions of other solutes owing to the electroneutrality constraint. Therefore, predictions for the nonreactive solutes, those that were modeled as not participating in any chemical reaction, will be discussed first. Then, solutes which are the products of the weathering reactions will be considered. Finally, the result of different deposition scenarios will be presented.

Nonreactive Solutes

The ionic loadings in the snowpack from field estimates are compared in table 3 with the measured export from discharge through the lake outlet during the snowmelt period. There are large differences between 1986 and 1987 both in the amount of precipitation and in its form. Rain inputs of water and solutes were negligible in 1986, but accounted for 18 percent of the water in 1987 and more than half the deposition of all solutes, except chloride. For the nonreactive solutes (sulfate and chloride in this model), the input should equal the output; clearly, this is not the case for sulfate and chloride during 1986 and 1987. The ionic deposition is calculated as the product of the ionic concentration and the volume of water, both of which are subject to measurement error.

Extensive snow surveys were performed to estimate the water volume of the snowpack. Techniques were refined and sampling intensity increased during 1986 and 1987 over the initial effort in 1985, as is evidenced by the good agreement in the water budget. Snowpack solute concentration was measured at eight points throughout the watershed, although not all sites were sampled for all events. Therefore, the discrepancy in the ionic budgets must be due either to sampling and analytical errors or to additional sources and mechanisms that regulate solute concentrations.

Sulfate input is less than one-half the sulfate output for 1986; yet the output exceeds the input for 1987. The difference between the form of the input (snow in 1986 and predominantly rain in 1987) may account for the difference, but there is insufficient data to make any conclusions. These budgets ignore input due to atmospheric dry deposition of sulfate and internal sources of sulfate. Contributions of sulfate by dry atmospheric deposition were evaluated using deposition to pine branches by Bytnerowicz and Olszyk (1988). An upper-bound estimate for the dry deposition of sulfur would result in only a 10 percent increase in the atmospheric inputs of table 3. A internal source of sulfate may exist. Volcanic glass was identified in the coarse silt fraction of soils at Emerald Lake (Wyles, 1986). Strong acid treatment of these soils resulted in sulfate release, which was attributed to the breakdown of vesicles that possibly contain sulfur dioxide. However, no quantitative estimate of this contribution to the sulfur budget has been made. Modifications to the model await further field observations of sulfur sources and regulating mechanisms.

The chloride budget is also variable, as might be expected, given the numerous opportunities for contamination during sample handling and the analytical precision at the low concentrations. The snow input number for 1986 may be somewhat high because there was a known chloride contamination from the plastic bags used for sampling that year. It was not possible to determine quantitatively the amount of contamination (M.W. Williams, Univ. of California, Santa Barbara, written commun., 1989). The chloride budget for 1987 would suggest, however, that the precision of these load estimates may be as low as ±50 percent if the variation in the chloride estimates arise primarily from sampling error and not from contamination or analytical imprecision.

Despite these differences in the budgets, the measured snowpack loadings for all solutes were used in the model. Although the differences on a seasonal basis appear to be large, the amount of error introduced in daily concentrations is only a few microequivalents per liter.
<table>
<thead>
<tr>
<th>Year</th>
<th>Water Volume</th>
<th>SBC</th>
<th>NH$_4^+$</th>
<th>NO$_3^-$</th>
<th>SO$_4^{2-}$</th>
<th>Cl$^-$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(10$^6$ m$^3$)</td>
<td>(10$^3$ eq)</td>
<td>(10$^3$ eq)</td>
<td>(10$^3$ eq)</td>
<td>(10$^3$ eq)</td>
<td>(10$^3$ eq)</td>
</tr>
<tr>
<td>1986</td>
<td>Snow input$^1$</td>
<td>2.4</td>
<td>7.5</td>
<td>2.9</td>
<td>4.1</td>
<td>3.4</td>
</tr>
<tr>
<td></td>
<td>Rain input$^2$</td>
<td>0.05</td>
<td>0.69</td>
<td>0.82</td>
<td>0.82</td>
<td>0.66</td>
</tr>
<tr>
<td></td>
<td>Total input</td>
<td>2.4</td>
<td>8.2</td>
<td>3.7</td>
<td>4.9</td>
<td>4.1</td>
</tr>
<tr>
<td></td>
<td>Output$^3$</td>
<td>2.3</td>
<td>63</td>
<td>0.1</td>
<td>9.1</td>
<td>9.9</td>
</tr>
<tr>
<td></td>
<td>Difference$^4$</td>
<td>0.1</td>
<td>-55</td>
<td>3.6</td>
<td>-4.2</td>
<td>-5.8</td>
</tr>
<tr>
<td></td>
<td>Ratio$^5$</td>
<td>.96</td>
<td>7.7</td>
<td>.027</td>
<td>1.9</td>
<td>2.4</td>
</tr>
<tr>
<td>1987</td>
<td>Snow input$^1$</td>
<td>0.80</td>
<td>3.2</td>
<td>3.0</td>
<td>3.4</td>
<td>2.3</td>
</tr>
<tr>
<td></td>
<td>Rain input$^2$</td>
<td>0.17</td>
<td>7.3</td>
<td>6.8</td>
<td>5.9</td>
<td>5.6</td>
</tr>
<tr>
<td></td>
<td>Total input</td>
<td>0.97</td>
<td>10.</td>
<td>9.8</td>
<td>9.3</td>
<td>7.9</td>
</tr>
<tr>
<td></td>
<td>Output$^3$</td>
<td>.75</td>
<td>34</td>
<td>0.01</td>
<td>6.3</td>
<td>5.2</td>
</tr>
<tr>
<td></td>
<td>Difference$^4$</td>
<td>0.22</td>
<td>-24</td>
<td>9.8</td>
<td>3.0</td>
<td>2.7</td>
</tr>
<tr>
<td></td>
<td>Ratio$^5$</td>
<td>0.80</td>
<td>3.4</td>
<td>.001</td>
<td>0.68</td>
<td>0.66</td>
</tr>
</tbody>
</table>

$^1$As determined from snowpack loadings at maximum accumulation (Dozier and others, 1989).

$^2$Includes rain and mixed rain and snow events that occurred during the melt period (Dozier and others, 1989) which ran from April 16, 1986 to September 30, 1986 and from March 25, 1987 to July 21, 1987.

$^3$As determined from mass export through lake outlet; water-quality observations were linearly interpolated to a daily basis.

$^4$Input less output; a positive value indicates net retention in the watershed and a negative value, a net release.

$^5$The ratio of output to input; values greater than one indicate that the watershed is a source of the solute and values less than one indicate that the watershed is a sink.
The results of the model simulations are presented in figures 8 and 9 for the combined inflows 1 and 2 and the lake outflow, respectively. Model results for streamwater quality cease when the model calculates that all the snow in the subbasin has melted; this occurred in mid July 1987. The “spikes” in concentration during the simulation period occur on during rain storms. Clearly these events were more important during 1987 than during 1986 as was also indicated by the budgets. The observations for the combined inflows (fig. 8) are an average of the measurements taken on the individual streams if more than one inflow was sampled on a particular day, or if more than one sample were collected during the day.

The upper three panels of each plot are for solutes that do not participate in the weathering reaction (Cl⁻, NO₃⁻, and SO₄²⁻). Three different elution scenarios are considered: an 80/20 release (80 percent of the solutes in the first 20 percent of the meltwater) with a maximum concentration factor (CF) of 8; a 40/20 release with a CF of 3; and a uniform release. For chloride, the 40/20 release rule seems to fit best; for nitrate, either the 40/20 for 1987 or 80/20 release rule for 1986 fits best. For sulfate, a uniform release fits the pattern best in 1986, although there is a systematic underprediction due to the net export of sulfate during this year. In 1987, the 40/20 rule appears to fit the initial stage of snowmelt best, yet during the rain events, the sulfate concentration does not respond as much as would be predicted by the model using either the 40/20 or 80/20 rule. The uniform release also does not describe the observations. In this case the sulfate added by the rain (twice as much as was in the snow, see table 3), continues to increase the stream concentration to levels above those observed. If there is a differential elution of sulfate (that is the 40/20 or 80/20 rule), then the stream sampling missed extreme changes in concentration. However, these concentrations would have to be much larger than those ever observed at Emerald Lake. Alternately, sulfate is not a nonreactive solute and is being retained in the watershed in 1987. Among the three solutes, sulfate exhibits the smallest range in concentration. This pattern with sulfate showing the least pronounced elution is opposite the preferential elution patterns observed by Brimblecombe and others (1987) where sulfate eluted first.

Putting aside the large rain-on-snow events in 1987 for the moment, consider 1986 and the April 1987 when an initial pulse from the snowpack would have been expected but did not occur. Two possible explanations can be advanced. One explanation is that solutes are eluted uniformly from the snowpack and that there are additional sources of nitrate and chloride, which account for the observed increases in concentration in the streamwater. Supporting this explanation is the observation that the peak nitrate concentrations observed in the inflow streams are roughly equivalent in 1986 and April, 1987 (approximately 12 µeq L⁻¹), yet the average concentration in the snowpack (assuming 10 percent and 100 percent biotic retention of nitrate and ammonium, respectively) is twice as large in 1987 (5 µeq L⁻¹) as in 1986 (2.5 µeq L⁻¹). The observed chloride pulse also may come from the soil solution where chloride from weathering or from the decomposition of organic matter may be concentrated during the winter (Peters, 1987).

A second explanation is that chloride and nitrate pulses arise from differential elution from the snowpack, but sulfate pulses from the pack are not observed in the stream owing to regulation by adsorption/desorption processes in the soils. Although soils cover only a small part of the watershed area, most flowpaths to the lake probably intersect the soils at some point. The constancy of the sulfate concentrations during the rain events in 1987 would also support sulfate regulation by the soils. However, such a regulating mechanism for sulfate has not been identified at Emerald Lake and would be unusual in such young soils.

The fundamental conclusion, regardless of the specific mechanism that accounts for the observed concentrations, is that the effective “acid pulse” to surface waters from this alpine snowpack is not as severe as would be predicted from laboratory studies or from point measurements as determined by snow lysimeters. For evaluation of the weathering model and the subsequent scenario analysis, the uniform and the 40/20 release rules will be considered.
Figure 8.— Observed and predicted concentrations of solutes at inflows 1 and 2.
Figure 9.—Observed and predicted concentrations of solutes in the lake outflow.
Weathering Products

The fourth through the seventh panels in figures 8 and 9 show the observed and the model predicted values for solutes affected by the weathering reactions (alkalinity, hydrogen ion, SBC, and silica). Two different solute release rules, uniform release and 40/20 release with a CF of 3, are considered. Model predictions are depicted as two lines that represent the upper and the lower bounds for predictions when the two different intercepts are used in the empirical weathering formulation as described above. As would be expected, the lines for the two different release rules cross each other during the melt. The 40/20 rule is lower at the beginning of the season for alkalinity because of the initial acid pulse, but ends higher than the uniform rule because the solutes already have been eluted, thus leaving less acidic water than the uniform elution case. Similarly, the 40/20 rule exhibits larger changes in concentration during rain events than the uniform release rule.

Considering the model predictions for the inflows during the calibration year 1986, the general pattern of dilution and recovery is reproduced for all solutes (fig. 8). Both silica and SBC are overpredicted during the latter part of the melt period by approximately 10 μeq L⁻¹; this overprediction could reflect biotic influences which are not included in the model such as diatomaceous uptake of silica and biotic uptake of calcium, magnesium and potassium during this period. Similarly, model predictions for the lake outflow (fig. 9) show the same general patterns, although SBC and silica are systematically underpredicted by 5 to 10 μeq L⁻¹ during the first three months of the simulation. This underprediction may result from errors in the distributed melt formulation or may indicate an additional source for these solutes such as ground water.

During the validation year 1987, however, the observed patterns of solute concentrations are not so well reproduced. The minimum alkalinity observed during snowmelt is correctly predicted, but the pattern of dilution during the initial phases of snowmelt is not captured correctly. In the inflows for the uniform release case (which better predicts alkalinity), silica is slightly overpredicted and SBC is substantially underpredicted. In addition, alkalinity is underpredicted during the early part of the snowmelt. These observations taken together suggest that cation exchange, not weathering, is supplying SBC and alkalinity (through consumption of hydrogen ion) during this first phase of melt. Cation exchange was not obvious during 1986 owing to the large volume of water that flowed through the system and to less intensive sampling of surface waters during the beginning of snowmelt. This additional alkalinity-generating mechanism would further ameliorate the acid pulse at the onset of snowmelt. Current model predictions overestimate the loss of alkalinity, and hence, provide a conservative prediction.
Deposition Scenarios

To assess the sensitivity of the watershed to different loadings of acidic solutes, two different atmospheric deposition scenarios were considered: one in which both sulfuric acid and ammonium nitrate loadings were doubled, and one in which both of these loadings were halved. Factor analysis indicates that the cations associated with sulfate and nitrate are hydrogen and ammonium, respectively (Lowell Ashbaugh, California Air Resources Board, written commun., 1988). Owing to the nitrogen cycle representation in ALF in which ammonium is exchanged for hydrogen ion, the scenarios are equivalent to changing the nitric acid loading to the system for this model. The results of these scenarios for the lake outlet are shown in figure 10. Again, each case is represented with a pair of lines bounding the predictions. The solid line is the half-loading case and the stippled line is the double-loading case. In figure 10, 1986 is indicative of a heavy snow year and 1987 represents a light snow year. The alkalinity of the system is not very sensitive to changes in loadings. The initial acid pulse causes the alkalinity in the outflow to decrease from about 20 μeq L^{-1} under current loadings (uniform release case in figure 9) to about 15 μeq L^{-1}. The acid pulse is more pronounced under the 40/20 release, as expected, with a minimum observed alkalinity of 8 μeq L^{-1} for 1986 conditions and 12 μeq L^{-1} for 1987 conditions as compared with minimum predicted alkalinities under current deposition of 10 and 15 μeq L^{-1} for 1986 and 1987, respectively. The snowpack loadings required to exhaust the alkalinity of the lake epilimnion, expressed in multiples of the current snow loading for 1986 and 1987, are summarized in table 4. No rain-on-snow events are assumed to occur.

<table>
<thead>
<tr>
<th>Table 4. Multiples of current deposition loadings of ammonium nitrate and sulfuric acid required to exhaust alkalinity of Emerald Lake</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Solute release rule</strong></td>
</tr>
<tr>
<td>Uniform release</td>
</tr>
<tr>
<td>40/20 release</td>
</tr>
<tr>
<td>80/20 release</td>
</tr>
</tbody>
</table>
Halving loading
Doubling loading

FIGURE 10—The effect of halving and doubling current nitrate and sulfate deposition on alkalinity of the lake outflow.
SUMMARY

This simple process-oriented model is able to capture the basic solute patterns during snowmelt in an alpine catchment where ground water is a minor contributor to streamflow. It includes an empirical representation of primary mineral weathering as the only alkalinity-generating mechanism. The model seems to be adequate during a heavy snow year, such as the one that served as the calibration year for this model. During a light snow year, there is evidence that cation exchange may be a substantial source of alkalinity during the initial phases of snowmelt. By not including cation exchange, the model overestimates the acidification during this period. The minimum observed alkalinity, which occurs through dilution, is predicted correctly by the model. The representation of the lake as simply a mixing volume with no additional chemical reactions is supported by the observations.

Model results indicate a change of 2 to 5 μeq L⁻¹ in the minimum alkalinity if the deposition of both ammonium nitrate and sulfuric acid is doubled. This estimate represents an upper bound on the alkalinity change because cation exchange is ignored. The predicted change in alkalinity would be insufficient to exhaust the alkalinity of the lake. Deposition would have to increase between two and 18 times the current loadings to accomplish that; the precise increment depends on hydrologic conditions and on the pattern of solute release from the snowpack. Although these increases in deposition seem to be very large, one should remember that current loadings to the system are low. The most likely scenario for acidification of Emerald Lake under conditions ranging from current to double the present-day deposition is an acidic rainstorm occurring during the latter part of snowmelt when the lake is at its minimum alkalinity owing to dilution from the meltwater. Such an event was observed in summer 1984. The infrequency of rainstorms in the Sierra Nevada, however, makes intensive sampling of storms difficult.

CONCLUSIONS

From this analysis of the data collected at Emerald Lake, additional avenues of research can be suggested both for modeling and field investigation. The simple form of ALF and its modest data requirements should allow for its widespread application in watersheds that are less intensively studied than Emerald Lake. The model can be calibrated with only water-quality and quantity measurements from the outflow of the lake. To run the model for any particular year requires an estimate of solute loadings in the pack. If there are no internal sources or sinks of sulfate and chloride in the watershed, these loadings can be estimated from the mass export from the watershed. Application of ALF to other watersheds in the Sierra Nevada would permit a regional assessment of the risk of episodic acidification.
The pattern of solute release from the snowpack is not consistent across the “nonreactive” solutes. In particular, sulfate, typically the mobile anion in such watersheds, has a surprisingly constant concentration in streamwater given the small amount of soil in the watershed. At a high altitude catchment in Scotland, sulfate concentration varied between 20 and 80 μeq/L during spring and summer, a factor of 4 (Cooper and others, 1987). Point measurements at Emerald Lake (Williams and Melack, 1988) and laboratory studies suggest that an acidic pulse should come from the pack at the beginning of snowmelt. However, stream concentrations of chloride, the least reactive solute, indicate a less intense ionic pulse than point measurements. Additional field observations and experiments need to be conducted to determine whether there are regulating mechanisms in the soil which control the sulfate concentration or whether these packs, due to the steep slope, ice lenses, mixing of waters at different stages of solute elution, or some other factor, do not in effect produce an acid pulse on the catchment scale. If the pack effectively elutes its solutes uniformly at this scale, an additional source for the observed nitrate and chloride peak must be identified. The apparent attenuation of the acid pulse, whatever its cause, renders this watershed much less sensitive to acidification from an acidified snowpack than if such a pulse existed.

The input-output budgets for sulfate and chloride were inconsistent between the two years of intensive study at Emerald Lake. The chloride input for 1986 is suspect due to sample contamination, but the sulfate budget deserves further study because the differences are too large to be attributable solely to sampling or analytical error. The most important distinction between the two years may be the size of the snowpack. The large amount of water in 1986 may have been able to leach more sulfate out of the watershed than in 1987, resulting in a net export in 1986 but a net retention in 1987. If this explanation is true, the chemical dynamics at Emerald Lake are more complicated than first thought both because sulfate is a reactive ion and because the flowpath through the watershed would determine the streamwater chemistry.

Paired mini-catchments having an area of several hundred square meters need to be identified at Emerald Lake to investigate sulfate regulation mechanisms. These catchments would represent the extremes in soil interaction; that is, one should have only bare bedrock and the other should be predominantly covered with soil. Meltwater draining the bare rock catchment would be affected by soil or talus, and thus, the existence or absence of an acidic pulse from the snowpack could be ascertained. In addition to runoff, soil solutions prior to and during the initial phases of melt need to be monitored to determine the effects of the soil on the “nonreactive” solutes.

Under current depositional loads and even with double the current loads, the Emerald Lake watershed is more susceptible to acidification from an acidic rainstorm which falls during the late melt period when the lake is its most dilute than to acidification from a acidic snowpack. The infrequency of such events in the Sierra Nevada makes sampling difficult. One field effort that is feasible and that may shed some light on the watershed’s response to acidic pulses is intensive sampling over the daily melt cycle. Sampling during early, mid and late melt season would provide information on the watershed’s chemical response to more acidic pulses, to clean pulses, and to clean pulses after the system has been strongly leached. In addition if sampling were carried out at points above and below the inflow meadow, then the influence of vegetation could be assessed.
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Dozier, Jeff, 1988, Topographic distribution of solar radiation over an alpine, snow-covered drainage basin [abs]: Eos Transactions of American Geophysical Union, v. 69, p. 1198.


Williams, M. W. and Melack, J.M., 1988, Snowpack runoff contributions to the major solute chemistry of lake and stream waters in a high-altitude watershed in the Sierra Nevada [abs], Eos Transactions of the American Geophysical Union, v. 69, p. 1198.

APPENDIX A.

Sample input/output from computer model

This attachment contains sample input which are necessary to run the computer model and samples of the output which the computer generates. The files are annotated by using Times-Roman typeface, the same typeface as this is written in. The contents of the files and any prompts which the computer types are written in Helvetica typeface, as these words are.

As the model runs, it executes a series of prompts for the names of the input files. Note that no output file name is requested since they are based on the name of the main input file. A sample dialog follows.

RUN DIST.86.CPL *User types the appropriate system command to execute the program

15 Mar 89 11:16:36 Wednesday
ENTER NAME OF MAIN INPUT FILE:
ILWAS>IN.WY86>1861/18A3.5 *User enters names of files are requested
ENTER NAME OF RAIN INPUT FILE:
ILWAS>IN.WY86>RAIN.WY86
ENTER NAME OF ICE ELEV INPUT FILE:
ILWAS>IN.WY86>ICELEV.WY86
ENTER NAME OF PRE 10 DAYS Q INPUT FILE:
ILWAS>IN.WY86>Q10DAY
ENTER NAME OF OBS STREAM INPUT FILE:
ILWAS>IN.WY86>IN.1
ENTER NAME OF OBS LAKE OUTPUT FILE:
ILWAS>IN.WY86>OUT.CONC
ENTER NAME OF OBS STREAM TEMP FILE:
ILWAS>IN.WY86>IN.TEMP
ENTER NAME OF OBS THERMOCLINE:
ILWAS>IN.WY86>THERMO.WY86
JD= 198 *Computer indicates which day it is working on
JD= 199
JD= 200

**** STOP
The required input files follow.

**Main Input File**

C
C
C
C These spaces are reserved for notes about the input
C *Lake initial cond: Julian day, ANC, CL, H, NH4, NO3, SBC, SIO2, SO4
C C
C
2  *kr flag => kr = F(Q) formula to use: 1 = actual, 2 = upper, 3 = lower
0.001 106 0.1 100  *partial pressure of CO2, % NO3 retained, day for Biotic influence
196 0.00 3.9 3.5 1.8 4.1 5.1 0.00 2.6  *ice chem initial condition
196 43.0 7.7 0.8 1 0.4 6.8 61.47 20.0 6.5  *epilimnion chem initial condition
196 37.0 3.6 1.6 6.1 19.7 58.88 20.0 6.6  *hypolimnion chem initial condition
198 365 202 365 1  *begin, end day of simulation, begin, end snowmelt hydrograph
198 2100.  *julian day and discharge
199 1800.
200 1600.

*Partial ionic loading of the snow pack in Equivalents: SBC, CL, NO3, NH4, SO4
8517.0 5619.7 4970.0 4107.0 9894.9
*Number of sub-units, total area, respective areas
7 1200000. 140298.82 132776.09 141176.47 396447.62 245115.45 134907.53 9278.03
*Sub-unit #, slope, aspect, latitude, elution (100 100 = uniform), concentration factor
1 33.7 42.3 36. 100 100 8.
2 30.6 27. 36. 100 100 8.
3 28.0 10. 36. 100 100 8.
4 33.8 305. 36. 100 100 8.
5 27.2 275. 36. 100 100 8.
6 33.7 253. 36. 100 100 8.
7 13.5 327. 36. 100 100 8.
*Representative values of inflow before snow melt.
*J-day, discharge, SBC, CL, NO3, NH4, SO4, H
198 2100. 8 1.5 1.99 1.07 0.77 0.94 1.73
199 1744.5 1.5 1.99 1.07 0.77 0.94 1.73
200 1551.3 1.5 1.99 1.07 0.77 0.94 1.73
201 1732.2 1.5 1.99 1.07 0.77 0.94 1.73

29
Rain data input file

297 14.8741 5.3597 22.9 24.94 20.2 14.79 *julian day, SBC CL NO3 NH4 SO4 H
324 24.2725 10.437 25.8 8.869 9.369 4.467
353 10.191 2.2567 2.9 4.435 2.498 2.951

Ice elevation file (elevation of ice/epilimnion interface):

196 4.5 *julian day, elevation
197 4.53125
198 4.5625

10-day average flow file (assumed to cover entire water year):

2763.2
2198.7
1915.1
1606.7
2170.1
1962
1791
1616.7
1715.2
2053.2

Chemical observations from main inflow:

198 36 8.77 8.04 7.34 1.4791 0.8 61.4 33.5 *julian day, ANC, Cl, NO3, SO4, H, NH4, SBC, SIO2
213 27.7 6.045 10.9675 7.6025 1.1977 0.325 47.5225 37.175
214 26.2 6.175 11.39 7.79 0.89149 0.15 48.69 36.35
217 26.8 4.795 9.79 6.66 1.2466 0 49.31 40.85

Chemical Observations for lake outflow:

213 5.98 1.0471E-06 0.2 32.5 5.8 10.15 8 45.65 56.4 *J-DAY,pH,H,NH4,ANC,CL,NO3,SO4,SIO2,SBC
214 5.98 1.0471E-06 0.1 35 5.4 9.9 7.2 45 53.2
218 6.04 9.1201E-07 0.1 35 6.9 10.6 6.6 43.9 56.7
Inflow stream temperature file:

198 0.13875 *Julian day and temperature
199 0.185
200 0.23125

Thermocline elevation file:

0.0 0.0 *elevation, Julian day. Code interpolates to obtain daily values.
0.0 25.0
7.6 72.0
8.5 99.
8.0 126.
2.5 191.
2.5 230.
0. 275.
0. 302.
5.5 317.
3.5 337.
1.0 351.
0. 357.
0. 365.

Output files There are fourteen output files produced by the program. All have a leading asterisk to allow for importation to a graphics package. Fields are tab delimited. The following is a list of these files with an explanation of what each contains.

D1861/18A3.5 sub-unit #1
D2861/18A3.5 sub-unit #2
D3861/18A3.5 sub-unit #3
D3861/18A3.5 sub-unit #4
D5861/18A3.5 sub-unit #5
D6861/18A3.5 sub-unit #6
D7861/18A3.5 sub unit #7
l861/18A3.5 snowmelt, calc inflow, observed inflow, for constructing bar graphs
L861/18A3.5 lake output
RL861/18A3.5 calculated vs observed, lake outflow
RS861/18A3.5 calculated vs observed, inflow
S861/18A3.5 combined stream inflow (d1 through d7)
V861/18A3.5 a text file for printing various daily information for visual review
VOL861/18A3.5 jday, total, ice, epilimnion, and hypolimnion volumes
#### D1861/18A3.5

<table>
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<tr>
<th>JDAY</th>
<th>QANCST</th>
<th>CLST</th>
<th>DOCST</th>
<th>HST</th>
<th>NH4ST</th>
<th>NO3ST</th>
<th>SBCST</th>
<th>SiO2ST</th>
<th>SO4ST</th>
<th>+/-BALI</th>
<th>PH</th>
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#### L861/18A3.5:

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<th>CL</th>
<th>DOC</th>
<th>H</th>
<th>NO3</th>
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<td>0.00</td>
<td>0.00</td>
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<td>1.99</td>
<td>-0.01</td>
<td>0.00</td>
<td>1.08</td>
<td>0.94</td>
<td>0.00</td>
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<tr>
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<td>0.00</td>
<td>0.00</td>
<td>0.56</td>
<td>0.00</td>
<td>0.00</td>
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<tr>
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<td>0.00</td>
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<td>0.00</td>
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<td>0.00</td>
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<tr>
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<td>36.00</td>
<td>8.77</td>
<td>-3.53</td>
<td>0.00</td>
<td>8.04</td>
<td>7.34</td>
<td>0.00</td>
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</table>

| CATIONSINPUT(S) | 199  | 1.50 | 0.00 | 0.00 | 0.00 | 1.74 | 0.00 | 0.00 | 0.77 |
| ANIONSCALC     | 199  | 0.00 | 1.99 | -0.01 | 0.00 | 1.08 | 0.94 | 0.00 |      |
| CATIONSCALC    | 199  | 40.82 | 0.00 | 0.00 | 0.56 | 0.00 | 0.00 | 0.00 |      |
| ANIONSCALC     | 199  | 0.00 | 36.93 | 1.99 | -0.01 | 0.00 | 0.97 | 0.94 | 0.00 |

#### L861/18A3.5:

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<th>DOCLK</th>
<th>HK</th>
<th>NH4L</th>
<th>NO3LK</th>
<th>SBCLK</th>
<th>SiO2LK</th>
<th>SO4LK</th>
<th>PHL</th>
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<td>0.3</td>
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<td>45.6</td>
<td>29.9</td>
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<td>-0.4</td>
<td>0.9</td>
<td>0.2</td>
<td>3.3</td>
<td>44.2</td>
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<td>0.2</td>
<td>2.8</td>
<td>41.5</td>
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<td>4.4</td>
</tr>
</tbody>
</table>

32
APPENDIX B.

Model Listing

The following is a listing of the program code. It has been written in standard FORTRAN-77. The only system-specific statements are the $INSERT statements, which are called $INCLUDE in some operating systems.

C
C MODIFIED FOR RAIN ON SNOWMELT.
C
C ADDED CALLS TO SUBROUTINES AFTER 150 CONTINUE
C
C
C
C C.... MAIN.F77
$INSERT COMMON/MAIN/
  COMMON/INIT/ RELO
  REAL*8 CN(9),SPCT(9),CROS(9)
  REAL*8 RS,RA,RL
  REAL*8 F1,F2,F3,F4,F5,F6,F7,F8,F9
  REAL*8 RELO
  REAL*8 QDIFF
  REAL*8 ANCAM,CLAM,DOCAM,HAM,NH4AM
  REAL*8 NO3AM,SBCAM,SI02AM,SO4AM
  INTEGER DATE
  CHARACTER FNAME*30,HEAD*80
  EQUIVALENCE (ANC,CN(1))
  EQUIVALENCE(ANCT,SPCT(1))
  TAB=CHAR(137)
C
C... Initial Values
C TO = REF. TEMP
C KHO = HENRYS K FOR CO2(g) AT TO
C R = UNIVERSAL GAS CONST
C RELO = INIT VALUE FOR % REL IONS IN FUNCTION RELFUNC
C
IRA1N1=0
RELO =0.00
R = 1.987
KHO=10.**(-1.47)
TO = 25.+273.16
INIT=0
IO=1
XAVE=0.0
DO 21 IA=1,1C
  QSUM(IA)=0.0
  CMELT(IA)=0.0
  CMELTO(IA)=0.0
  CREL(IA)=0.0
  CRELO(IA)=0.0
  IOFF(IA)=1
  IOFF2(IA)=1
  IOFF3(IA)=1
  IOFF4(IA)=1
21 CONTINUE
C... Integer Lake/Stream Observed Day (ILOD,ISOD) COUNTERS USED WRITING
C TO A FILE WHICH CONTAINS OBSERVED DATA VS. CALCULATED DATA (R^2)
ILOD=1
ISOD=1
C...IR IS A DAILY COUNTER FOR RAIN DAY CONCENTRATIONS
IR=1
C
C... CALL FILER TO OPEN IN/OUTPUT FILES, READ INITIAL CONDITIONS
CALL FILER
C... CALL LAKE AND GIVE IT THESE CONDITIONS!
CALL LAKE(1)
C... CALL DISTM AND DISTRIBUT THE DISCHARGE AND IONS, CALC RELEASE RULE
CALL DISTM
C
C... BEGIN DAILY TIME STEPS
IPC=1
NPTS=JDN-JD1+1
DO 50 I=JD1,JDN,1
NRR=0
XDAT(IPC)=FLOAT(I)
JDAY=I
WRITE(*,*)' JDAY',JDAY
WRITE(*,*)QSU(4,JDAY)
C
C...READ DAILY INFLOW TEMP IN CELCIUS
READ(20,*)JY,ST
C
C...CONVERT INFLOW TEMP TO KELVINS
ST=ST+273.16
C
C...MEMORY, FOR Q vs. Kr
IF(MEM.EQ.1)THEN
QT=0.0
DO 52 J=0,9
52 QT=QT+SH(I-J)
QT=QT/10.
ELSE
QT=SH(I)
ENDIF
C
C... ARE WE BEFORE, DURING, OR AFTER THE MELT.
IF(JDAY.LT.JBM)THEN
GOTO 200
ELSEIF(JDAY.GT.JEM)THEN
GOTO 300
ENDIF
C
C**************************************
C... HERE WE DO DISTRIBUTED MELT, JDAY .GE. JBM .AND. JDAY .LT. JEM
C... THE Q FOR EACH SUB UNIT IS IN ARRAY AQSU(IA,ND) WHERE IA IS THE
C... SUB UNIT AND ND IS THE DAY. ANOTHER ARRAY, ND(IA) HAS THE DAY
C... THAT EACH SUB UNIT DRIES UP
C
C... FIRST WE CHECK FOR RAIN ON SNOW.
C
IF(RR(I).GT.0.0)THEN
SBC=SBCR(IR-1)
CL=CLR(IR-1)
NO3=NO3R(IR-1)
NH4=NH4R(IR-1)
SO4=S04R(IR-1)
H=HR(IR-1)
DOC=DOCR(IR-1)
IF(DOC.LT.0.0)THEN
DOC=0.0
ENDIF
WRITE(*,*)' WARNING: NEGATIVE DOC ON RAIN JDAY =',JDAY
ENDIF
C
WRITE(*,*)'CALL NRWEATH 1'
CALL NRWEATH
DO 152 IJK=1,9
CROS(IJK)=CN(IJK)
152 CONTINUE
ENDIF
C
C... CALCULATE PRECENTAGE OF IONS RELEASED FROM EACH SUB UNIT
C... CONVERT EQ/M**3 TO uEQ/L, WHERE QSU IS IN METERS CUBED
DO 150 IA=1,NSU
IF(JDAY.GT.ND(IA,1))GOTO 150
DO 151 IC=1,9
IF (IC.EQ.1.OR.IC.EQ.8) THEN
CN(IC)=0.0
GOTO 151
ELSE
T=TIME(JDAY,IA,IC)
IF(JDAY.GT.TSTAR(JDAY,IA,IC)) THEN
C.. UNITS UEQ/L
C IF(IC.EQ.2.AND.IA.EQ.4)WRITE(*,*)'AFTER T*
CN(IC)=AB(JDAY,IA,IC,3)*EXP(-AB(JDAY,IA,IC,4)*T)*1000.
ELSE
C IF(IC.EQ.2.AND.IA.EQ.4)WRITE(*,*)'BEFORE T*
CN(IC)=AB(JDAY,IA,IC,1)*EXP(-AB(JDAY,IA,IC,2)*T)*1000.
ENDIF
C IF(IC.EQ.9.AND.IA.EQ.4)WRITE(*,*)AB(JDAY,IA,IC,1),
C & AB(JDAY,IA,IC,2),
C & AB(JDAY,IA,IC,3),
C & TSTAR(JDAY,IA,IC)
151 CONTINUE
IF(IA.EQ.4) THEN
C PRINT OUT MELT WATER, BEFORE ROCK REACTION
C WRITE(*,33)JDAY,TAB,ANC,TAB,CL,TAB,DOC,TAB,H,
ENDIF
C... CALC HYDROGEN ST THE ELUDED IONS HAVE CHARGE BALANCE
C H=CL+NO3+SO4-SBC-NH4
C... SEND VALUES IN UEQ/L
C DOC=FD(H,SBC,NH4,SO4,CL,NO3)
IF(DOC.LT.0.0) THEN
WRITE(*,*)' WARNING: NEGATIVE DOC ON JDAY =',JDAY
WRITE(*,*)' DOC = ',DOC
DOC=0.0
ENDIF
IF(ANC.LT.0.01)ANC=0.0
IF(SIO2.LT.0.01)SIO2=0.0
C
C... THESE VALUES ARE FOR AFTER MELT (AM) WHEN INFLOW CONC TO LAKE IS CONST.
C THESE VALUES ARE FROM THE LAST CONTRIBUTING SUBUNIT, IF TWO SUBUNITS
C DRY UP ON THE SAME DAY, THE THE ONE INPUTED LAST IS USED.
C
ANCAM=ANC
CLAM=CL
DOCAM=DOC
HAM=H
C
C     WRITE(*,*)' IA = ', IA
C     WRITE(*,*)' ANC = ', ANC
C     WRITE(*,*)' CL = ', CL
C     WRITE(*,*)' DOC = ', DOC
C     WRITE(*,*)' H = ', H
C     WRITE(*,*)' NH4 = ', NH4
C     WRITE(*,*)' NO3 = ', NO3
C     WRITE(*,*)' SBC = ', SBC
C     WRITE(*,*)' SI02 = ', SI02
C     WRITE(*,*)' S04 = ', S04
C     IF(IA.EQ.4) THEN
C         WRITE(*,*)'
C     ENDIF
C     CALL NRWEATH
C     IF(IA.EQ.4) THEN
C         WRITE(*,*)' CL = ', CL
C     ENDIF
C     WRITE(*,*)' AFTER CALL TO NRWEATH
C     WRITE(*,*)' ANC = ', ANC
C     WRITE(*,*)' CL = ', CL
C     WRITE(*,*)' DOC = ', DOC
C     WRITE(*,*)' H = ', H
C     WRITE(*,*)' NH4 = ', NH4
C     WRITE(*,*)' NO3 = ', NO3
C     WRITE(*,*)' SBC = ', SBC
C     WRITE(*,*)' SI02 = ', SI02
C     WRITE(*,*)' S04 = ', S04
C     WRITE(*,*)'
C
C... HERE WE TRACE THE OUTFLOW OF EACH SUB UNIT, CHECK FOR RAIN FIRST
IF(RR(JDAY).GT.0.0) THEN
    XCENT=AREA(IA)/AREAT
    DO 169 IC=1,9
        CN(IC) = ((CN(IC)*QSU(IA,JDAY) ) +
                (CROS(IC)*XCENT*RR(JDAY)) )/
                (QSUUA, JDAY) +RR(JDAY) *XCENT)
    169 CONTINUE
    CALL NROPEN
ENDIF
PH=-LOG10(H/1000000)
WRITE (28+IA,43)JDAY,TAB, QSU(IA, JDAY) , TAB,ANC,TAB,CL, TAB, DOC, 
& TAB, H, TAB, NH4, TAB, NO3, TAB, SBC, TAB, SI02, TAB, S04, TAB, CB, TAB, PH
C
DO 170 IC=1,9
    SPCA(IA,IC)=CN(IC)
170 CONTINUE
150 CONTINUE
C
C... MIX ALL SUB UNIT OUTPUT BEFORE MIXING INTO LAKE
DO 180 IC=1,9
    CN(IC)=0.0
    QCSUM=0.0
DO 185 IA=1,NSU
IF(JDAY.GT.ND(IA,1))THEN
  IF(RR(JDAY).GT.0.0)THEN
    XCENT=AREA(IA)/AREAT
    CN(IC)=(CROS(IC)*RR(JDAY)*XCENT+QCSUM*CN(IC))
    / (QCSUM+RR(JDAY)*XCENT)
  ENDIF
ELSE
  CN(IC)=(SPCA(IA,IC)*QSU(IA,JDAY)+QCSUM*CN(IC))
  / (QCSUM+QSU(IA,JDAY))
ENDIF
CONTINUE
CONTINUE

C... TEST FOR LOW ANC AND PRINT OUT PH
C... UNITS ARE MICRO EQUIVALENTS
C
PH=-LOG10(H*1000000.0)
C
IF(ABS(ANC-H).LT.0.5)THEN
  WRITE(*,*) 'ANC',ANC
  WRITE(*,*) 'H',H
ENDIF

C
C... WRITE COMBINED STREAMS OUTPUT TO FILE 13
C
PH=-LOG10(H/1000000)
WRITE(13,43)JDAY,TAB,QI(JDAY),TAB,ANC,TAB,CL,TAB,DOC,TAB,H,
&TAB,NH4,TAB,NO3,TAB,SBC,TAB,SIO2,TAB,S04,TAB,PH
ENDIF
43 FORMAT(2X,'ANC',4X,'CL',3X,'DOC',4X,'H',4X,
&'NH4',3X,'NO3',3X,'SBC',2X,'SIO2',3X,'S04')
3 FORMAT(13(F5.1,IX),/)
C
C... IF STREAM OBSERVATIONS EXIST FOR THIS DAY, WRITE OUT CALC AND OBS DATA
C FOR COMPARISON (R^2, TAB DELIMITED, OUTPUT FILE)
C
IF(JDAY.EQ.OSJD(ISOD))THEN
  OSH(ISOD)=OSH(ISOD) /1000000
  OSPH=-LOG10(OSH(ISOD))
  OSH(ISOD)=OSH(ISOD)*1000000
  WRITE(7,43)JDAY,TAB,QI(JDAY),TAB,ANC,TAB,CL,TAB,DOC,TAB,H,
  &TAB,NH4,TAB,NO3,TAB,SBC,TAB,SIO2,TAB,S04,TAB,PH
  &TAB,OSANC(ISOD),TAB,OSCL(ISOD),TAB,OSH(ISOD),TAB,
  &OSNH4(ISOD),TAB,OSNO3(ISOD),TAB,OSBBC(ISOD),TAB,OSO2(ISOD)
  &TAB,OSPH(ISOD)
  ISOD=ISOD+1
ENDIF
4 FORMAT(13(F5.1,IX),/)
C
C... BEGIN LAKE MIXING
GOTO 500
C
C... HERE WE TAKE CONCENTRATION FROM INPUT PREVIOUS TO MELT
C
200 CONTINUE

C... SNOW MELT DATA: jday, Q(M**3), SBC(uEQ/L), CL, NO3, NH4, SO4
READ (9, *) DAY, QI(I), SBC, CL, NO3, NH4, SO4, H

C WRITE(*,*)'CALL RAIN 1'
CALL RAIN
C WRITE(*,*)'CALL NRWEATH 3'
CALL NRWEATH

PH = -LOG10(H/1000000)
WRITE(13, 43) JDAY, TAB, QI(JDAY), TAB, ANC, TAB, CL, TAB, DOC, TAB, H,
& TAB, NH4, TAB, NO3, TAB, SBC, TAB, SIO2, TAB, SO4, TAB, CB, TAB, PH

C... IF STREAM OBSERVATIONS EXIST FOR THIS DAY, WRITE OUT CALC AND OBS DATA
C FOR COMPARISON (R2, TAB DELIMITED, OUTPUT FILE)

C IF(JDAY.EQ.OSJD(ISOD)) THEN
OSH(ISOD) = OSH(ISOD) / 1000000
OSH(ISOD) = OSH(ISOD) * 1000000
WRITE(7, 43) JDAY, TAB, QI(JDAY), TAB, ANC, TAB, CL, TAB, H,
& TAB, NH4, TAB, NO3, TAB, SBC, TAB, SIO2, TAB, SO4, TAB, PH,
& TAB, OSANC(ISOD), TAB, OSCL(ISOD), TAB, OSH(ISOD), TAB,
& OSNH4(ISOD), TAB, OSNO3(ISOD), TAB, OSBC(ISOD), TAB, OSI02(ISOD),
& TAB, OSS04(ISOD), TAB, OSPH
ISOD = ISOD + 1
ENDIF
GOTO 500

300 CONTINUE
C
C... THESE VALUES ARE THE LAST VALUES OF SNOW MELT
C... AND ARE USED TO DRIVE THE MODEL WHEN ALL SUB UNITS
C... HAVE DRIED UP.
ANC = ANCAM
CL = CLAM
DOC = DOCAM
H = HAM
NH4 = NH4AM
NO3 = NO3AM
SBC = SBCAM
SIO2 = SIO2AM
SO4 = SO4AM

400 CONTINUE
C
C
C... EQUILIBRATE USING NEWTON-RAPHSON METHOD

C WRITE(*,*)'CALL RAIN 2'
CALL RAIN
C WRITE(*,*)'CALL NRWEATH 4'
CALL NRWEATH

PH = -LOG10(H/1000000)
WRITE(13, 43) JDAY, TAB, QI(JDAY), TAB, ANC, TAB, CL, TAB, DOC, TAB, H,
& TAB, NH4, TAB, NO3, TAB, SBC, TAB, SIO2, TAB, SO4, TAB, CB, TAB, PH
C
C... IF STREAM OBSERVATIONS EXIST FOR THIS DAY, WRITE OUT CALC AND OBS DATA
C FOR COMPARISON (R2, TAB DELIMITED, OUTPUT FILE)
IF (JDAY.EQ.OSJD(ISOD)) THEN
  OSH(ISOD) = OSH(ISOD)/1000000
  OSPH = -LOG10(OSH(ISOD))
  OSH(ISOD) = OSH(ISOD)*1000000
  WRITE(7,43) JDAY, TAB, QI(JDAY), TAB, ANC, TAB, CL, TAB, H
  & , TAB, NH4, TAB, NO3, TAB, SBC, TAB, SIO2, TAB, SO4, TAB, PH
  & , TAB, OSANC(ISOD), TAB, OSCL(ISOD), TAB, OSH(ISOD), TAB,
  & OSNH4(ISOD), TAB, OSNO3(ISOD), TAB, OSSBC(ISOD), TAB, OSSO4(ISOD)
  & , TAB, OSI02(ISOD), TAB, OSPH
  ISOD = ISOD + 1
ENDIF

C  C... HERE WE JUMP BACK IN AFTER THE DISTRIBUTED MELT, MELT ENDED
C... AND BEFORE THE BEGINNING OF MELT

500 CONTINUE

C  C... OUTPUT IONIC STRENGTHS OF INPUT(SNOW MELT OR RAIN),
C... CALCULATED LAKE INFLOW, AND OBS STREAM INFLOW.
C
C  WRITE(*,*)' CALL IONIC 1'
  CALL IONIC

C  C... MIX WITH LAKE VOLUME
C  WRITE(*,*)' CALL LAKE 1'
  CALL LAKE(0)

C  C... PRINT OUTPUT
  DO 40 J=1,9
    CN(J) = QI(JDAY) * CN(J) * 0.001
  WRITE(15,*)' OUTFLOW LOADING, EQ'
  WRITE(15,4)
  WRITE(15,7)(CN(IJI), IJI=1,9)
  7 FORMAT (13(F8.1, IX) , /)
  88 FORMAT (13(F8.2, A2))
  IPC = IPC + 1
  IRAIN1 = 0
40 CONTINUE

C  C
C  C  CLOSING FILES
  CLOSE(6)
  CLOSE(18)
  CLOSE(17)
  CLOSE(8)
  CLOSE(9)
  CLOSE(10)
  CLOSE(11)
  CLOSE(12)
  CLOSE(13)
  CLOSE(14)
  C  CLOSE(15)
  CLOSE(5)
  CLOSE(7)
  CLOSE(19)
  CLOSE(20)

C*** USED TO CHECK LAKE MIXING
  CLOSE(29)
  CLOSE(30)
  CLOSE(31)
  CLOSE(32)
  CLOSE(33)
  CLOSE(34)
DOUBLE PRECISION FUNCTION RELFUNC(XM, IA, MS)
  REAL*8 XM

C... THE EQUATIONS
C... DEPENDING ON THE VALUE OF IA, A SUB UNITS HAS AN RELEASE EQUATION
C... THE XM IS CUMILATIVE MELT (%), AND REL IS CUMULATIVE RELEASE (%)
C
IF(MS.EQ.11) THEN
  RELFUNC = XM
ELSEIF(MS.EQ.82) THEN
  IF(XM.LE.0.2) THEN
    RELFUNC = 4.DO*XM
  ELSE
    RELFUNC = 0.25DO*XM + 0.75DO
  ENDIF
C WRITE(*,*) 'RELFUNC 82 =', RELFUNC
ENDIF
RETURN
END

SUBROUTINE DISTM
SINSERT COMMON/MAIN/
  REAL*8 CN(9), SPCT(9)
  REAL*8 TMASS, PMASS, PWAT
  EQUIVALENCE (ANCT, SPCT(1))
C
C HERE WE DIVIDE MASS OF SNOW BY AREA, CALCULATE DAILY RADIATION
C DAILY DISCHARGE FROM EACH SUB UNIT, COEFFICENTS FOR THE RELEASE
C RULE FOR IONS, A1,A2,B1,B2,NDAY
C
C.. DIVIDE TOTAL MASS OF SNOW AMONG SUB UNITS BASED ON
C.. RELATIVE AREAS
DO 66 IA = 1, NSU
  QSUTOT(IA) = (AREA(IA)/AREAT)*QTOT
C WRITE(*,*) 'DISTM QSUTOT(', IA, ')=', QSUTOT(IA)
DO 66 IC = 1, 9
  IF(IC.EQ.1 OR IC.EQ.8) GOTO 66
  SPCAT(IA, IC) = (AREA(IA)/AREAT)*SPCT(IC)
66 CONTINUE
C WRITE(*,*) 'SPCAT(', IA, ', IC, ')=', SPCAT(IA, IC)
C
C.. DAILY LOOP FOR SOLAR RADITATION AND DISCHARGE
C
DO 100 ID = JBM, JEM
C WRITE(*,*) 'JDAY', ID
C WRITE(*,*) 'MAIN QC, ID, ') = ', QS
C
C.. CALCULATE PERCENTAGE INSOLATION ON SUB UNITS PER UNIT AREA
  SRAD = 0.0
DO 110 IA = 1, NSU
  IF(IOFF(IA).EQ.0) GOTO 110
  RSB = RSSLOPE(IA)
  RA = RASPECT(IA)
  RL = RLAT(IA)
  CALL SOLRAD(ID, RSB, RA, RL, RAD)
  SRAD = SRAD + RAD
110 CONTINUE
C
40
RSU(IA) = RAD

C 110 CONTINUE
C
C... CALCULATE AREA INTO MELT
DEM=0.0
DO 115 IA=1,NSU
  IF(IOFF(IA).EQ.0)GOTO 115
  D=AREA(IA)*RSU(IA)/SRAD
  DEM=DEM+D
115 CONTINUE
C
C... CALCULATE THE PRECENT DISCHARGE FOR EACH SUBUNIT
DO 120 IA=1,NSU
  IF(IOFF(IA).EQ.0)GOTO 120
  RCENT = (AREA(IA)*RSU(IA)/SRAD)/DEM
  QSU(IA,ID)=RCENT*QS
120 CONTINUE
C
C... CHECK IF SUB UNIT CUMULATIVE DISCHARGE HAS EXCEEDED TOTAL WATER
WRITE(*,*) ' '  IRT=1
DO 130 IA=1,NSU
  WRITE(*,*)' QSUM(', IA, ')=',QSUM(IA)
  IF(IOFF(IA).EQ.0)GOTO 130
  TQSUM(IA) = QSUM(IA) + QSU(IA,ID)
  WRITE(*,*)' MAIN TQSUM(', IA, ')=',TQSUM(IA)
  WRITE(*,*)' QSUTOT(', IA, ')=QSU(', IA, ',ID)
  QSU(IA,ID)= QSUTOT(IA)-QSUM(IA)
  WRITE(*,*)' DRIED UP NEW QSU(', IA, ',ID)=',QSU(IA,ID)
  QS =QS - QSU(IA,ID)
  IRT = 0
ENDIF
IF(IOFF(IA).NE.0)THEN
  ND(IA,1)=ID
ENDIF
130 CONTINUE
C
C... RECALCUATE PRECENT INSOLATION PER SUB UNIT IF A UNIT 'DRYS UP'
IF(IRT.EQ.0)GOTO 5
C
C... UPDATE THE SUM OF WATER LEAVING A SUB UNIT
DO 140 IA=1, NSU
  IF(IOFF2(IA).EQ.0)GOTO 140
  QSUM(IA)=QSUM(IA)+QSU(IA,ID)
WRITE(*,*)' MAIN QSUM(', IA, ')=',QSUM(IA)
IF(IOFF(IA).EQ.0)IOFF2(IA)=0
140 CONTINUE
100 CONTINUE
C
DO 131 IA=1, NSU
C131 WRITE(*,*)' ND(', IA, ',1)=',ND(IA,1)
C DO 301 I=JBM,JEM
  WRITE(*,*) 'QSU(', I, ')=',QSU(I)
C
C... CALC THE PARAMETERS FOR CONCENTRATION
C
DO 300 IA=1, NSU
C
DO 250 IC=1,9
C WRITE(*,*)' IA=',IA,' IC=',IC
K1=1
JBS=JBM
450 CONTINUE
C... NO ANC AND SIO2 CONTRIBUTED, DOC IS CALCULATED LATER
IF(1C.EQ.1.OR.1C.EQ.8.OR.1C.EQ.3)GOTO 250
PMASS=DBLE(MSEL(IA,1))/100.
PWAT=DBLE(MSEL(IA,2))/100.
TMASS=SPCAT(IA,1C)
NQ=ND(IA,1)
CFS=CF(IA)
CALL ELUTR(QSU,JBS,NQ,IA,TMASS,PWAT,PMASS,CFS,A1,B1,A2,B2,NDAY)
ND(IA,2)=NDAY
C WRITE(*,43)JBS,TAB,IA,TAB,1C,TAB,Al,TAB,Bl,TAB
C & ,A2,TAB,B2,TAB,SPCAT(IA,1C) ,TAB,ND (IA,2)
43 FORMAT(13,Al,II,Al,11,5(A1,E12.6E2),Al,13)
T=1
IF(MSEL(IA,1).EQ.MSEL(IA,2))THEN
QSUM(IA)=0.0
DO 399 I=JBS,JEM
QSUM(IA)=QSUM(IA)+QSU(IA,I)
399 CONTINUE
ENDIF
DO 400 I=JBS,JEM,1
AB(I,IA,IC,1)=A1
AB(I,IA,IC,2)=B1
AB(I,IA,IC,3)=A2
AB(I,IA,IC,4)=B2
IF(MSEL(IA,1).EQ.MSEL(IA,2))THEN
AB(I,IA,IC,1)=TMASS/QSUM(IA)
AB(I,IA,IC,2)=0.0
AB(I,IA,IC,3)=TMASS/QSUM(IA)
AB(I,IA,IC,4)=0.0
ENDIF
IF(I.GT.ND(IA,2))THEN
CN(IC)=AB(I,IA,IC,3)*EXP(-AB(I,IA,IC,4)*T)
ELSE
CN(IC)=AB(I,IA,IC,1)*EXP(-AB(I,IA,IC,2)*T)
ENDIF
TIME(I,IA,IC)=T
TSTAR(I,IA,IC)=ND(IA,2)
IF(I.EQ.ND(IA,2))T=0
T=T+1
SPCAT(IA,IC)=SPCAT(IA,IC)-CN(IC)*QSU(IA,I)
XGANG=CN(IC)*QSU(IA,I)
C WRITE(*,57)I,TAB,SPCAT(IA,IC) ,TAB,XGANG
57 FORMAT(13,Al,E12.6E2,Al,E12.6E2)
IF(I.EQ.JDR(KI))GOTO 410
400 CONTINUE
GOTO 250
410 IF(I.EQ.2)THEN
BANG=CLR(KI)
ELSEIF(I.EQ.4)THEN
BANG=HR(KI)
ELSEIF(I.EQ.5)THEN
BANG=NH4R(KI)
ELSEIF(I.EQ.6)THEN
BANG=N03R(KI)
ELSEIF(I.EQ.7)THEN
BANG=SBCR(KI)
ELSEIF(I.EQ.9)THEN
BANG=S04R(KI)
ELSE
BANG=CLR(KI)
ENDIF
WRITE(*,*)BANG
ENDIF
C... ADD RAIN STORAGE * RAIN CONC TO SNOW PACK ON THIS RAIN DAY
BANG=RS(I)*(AREA(A)/AREAT)*BANG/1000.
C
WRITE(*,*)BANG
C
SPCAT(IA,IC)=SPCAT(IA,IC)+BANG
C
WRITE(*,*)SPCAT(IA,IC)
JBS=JDR(KI)+1
C
WRITE(*,*)'RAIN DAY =',JDR(KI)
KI=KI+1
IF(JBS.GT.ND(IA,1))THEN
GOTO 250
ELSE
GOTO 450
ENDIF
250 CONTINUE
300 CONTINUE
C
RETURN
END
C
SUBROUTINE ELUTR(Q,JBM,NO,IA,TMASS,PWAT,PMASS,CF,Al,Bl,A2,B2,NDAY)
IMPLICIT DOUBLE PRECISION(A-H,O-Z)
DIMENSION Q(10,1000)
REAL*8 MPR,MPR?
C
This program takes an elution rule (e.g., 80% of the ions in 20% of
the melt, and the concentration on a given day and returns the two
parameters for an exponentially declining concentration. A simple
Newton's method of solution is used.
C
TWAT=0
DO 10 I=JBM,NQ
TWAT=TWAT+Q(IA,I)
10 CONTINUE
C
WRITE(*,*)'TWAT-',TWAT,' JBM=',JBM,' NQ-',NQ
SUMQ=0
DO 20 I=JBM,NQ
SUMQ=SUMQ+Q(IA,I)
IF(SUMQ.LT.PWAT*TWAT) THEN
GOTO 20
ELSE
GOTO 30
ENDIF
20 CONTINUE
C
WRITE(*,*)'ERROR IN CALC. OF DAYS REQD FOR FLOW.'
STOP
C
NDAY=I
C
WRITE(*,*)'NDAY= ',NDAY
C
ready to begin Newton's method solution; MAKE INITIAL GUESS
B1=.03
CAVE=TMASS/TWAT
MPR=TMASS*PMASS
C0=CF*CAVE
50 XSUM=0.0
FBR=0.0
J=JBM
CK=NDAY-JBM+1
DO 100 I=1,CK
CN = C0*DEXP(-B1*DBLE(I))
XSUM=XSUM + CN*Q(IA,J)
FBR =FBR + DBLE(I)*Q(IA,J)*C0*DEXP(-B1*DBLE(I))
J=J+1
100 CONTINUE
F=MPR-XSUM
C=F-FBR*B1
IF(DABS(F).GT.1.D-06)THEN
   B1=-C/FBR
   GOTO 50
ENDIF
A1=CF*CAVE
C********
CO = CN
MPR = TMASS -MPR
B2 = .025
JK=J
150 J=JK
XSUM=0.0
FBR=0.0
DO 110 I=1,NQ-NDAY
   CN = Q(I,J)*CO*DEXP(-B2*DBLE(I))
   XSUM=XSUM+CN
   FBR =FBR + DBLE (I)-*Q(I,J)*CO*DEXP (-B2*DBLE(I))
   J=J+1
110 CONTINUE
F=MPR-XSUM
C=F-FBR*B2
IF(DABS(F).GT.1.D-06)THEN
   B2=-C/FBR
   GOTO 150
ENDIF
A2=CO
RETURN
END

SUBROUTINE SOLRAD(JDAY,I,A,LO,R4)
COMMON/RAD/ JD,D
REAL*8 D
REAL*8 I,LO,A,L1,L2,X,Y,R1,E,T
REAL*8 T6,T7,T1,T0,T2,T3,D1,R4
INTEGER JDAY
C
C... THIS IS FROM SWIFT 1976, TO CALCULATE POTENTIAL SOLAR RADIATION.
C
C... I,LO,A, ARE CONVERTATED FROM DEGREES TO RADIANS
I=DR(I)
LO=DR(LO)
A=DR(A)
C
X= DCOS(I) *DSIN(LO) + DSIN(I) *DCOS(LO) *DCOS(A)
L1=DATAN(X)
X= DSIN(I) *DSIN(A)
Y= DCOS(I) *DCOS(LO) - DSIN(I) *DSIN(LO) *DCOS(A)
L2=DATAN(X/Y)
C
L1=RD(L1)
L2=RD(L2)
C...
L1 AND L2 ARE NOW IN DEGREES
C
WRITE(*,*)'L1 =',L1
C
WRITE(*,*)'L2 =',L2
C...
CONVERT JDAY BEGINNING OCT 1, TO JULIAN DAY BEGINNING JAN 1
IF(JDAY.LE.92)THEN
   JD=JDAY+273
ELSEIF(JDAY.GT.92)THEN
   JD=JDAY-92
ENDIF
C
D = \text{FUNC1}(0.400, 23.300, 10.000)

WRITE(*, *) ' D (DEG) = ', D

C ... CONVERT LO FROM RADIANS TO DEGREES
LO = \text{RD}(LO)

E = \text{FUNC1}(1.000, 0.01670, -3.000)
R1 = 60.0000 * 1.95 / (E^2)

WRITE(*, *) ' R1 = ', R1

T = \text{FUNC2}(L1)
T7 = T - L2
T6 = -T - L2
T = \text{FUNC2}(LO)
T1 = T
T0 = -T
IF(T7.LT.T1) THEN
  T3 = T7
ELSE
  T3 = T1
ENDIF
IF(T6.GT.T0) THEN
  T2 = T6
ELSE
  T2 = T0
ENDIF

WRITE(*, *) ' L2, L1, T3, T2 ', L2, L1, T3, T2

RETURN  
END

\begin{verbatim}
DOUBLE PRECISION FUNCTION FUNC3(V, W, X, Y, R1)
COMM)
COMM.
COMMON/RAD/ JD, D
REAL*8 D
REAL*8 V, W, X, Y, PI, R1
REAL*8 VV, WW, XX, YY, A, B, DD
PI = 3.141592654DO
DD = DR(D)
WW = DR(W)
XX = DR(X)
YY = DR(Y)
VV = DR(V)
A = DSIN(DD) * DSIN(WW) * (X - Y) / 15.0DO
B = DDCOS(DD) * COS(WW) * (DSIN(XX + VV) - DSIN(YY + VV)) * 12.0DO / PI
WRITE(*, *) ' R1 = ', R1
RETURN
END

DOUBLE PRECISION FUNCTION FUNC2(Y)
COMM.
COMMON/RAD/ JD, D
REAL*8 D
REAL*8 X, DD, YY, R1, Y
DD = DR(D)
YY = DR(Y)
X = (-DTAN(YY) * DITAN(DD))
IF(X.GT.1.000) THEN
  WRITE(*, *) ' SOLRAD X = ', X
  X = 1.0DO
ENDIF
IF(X.LT.-1.000) THEN
  WRITE(*, *) ' SOLRAD X = ', X
  X = -1.0DO
\end{verbatim}
DOUBLE PRECISION FUNCTION FUNC1(W,X,Y)

ESTIMATE DECLINATION, RADIUS VECTOR OF SUN, FUNC OF JULIAN DAY

COMMON/RAD/ JD,D
REAL*8 D
REAL*8 W,X,Y,V,R1
V=(DBLE(JD)+Y)*0.986DO
V=DR(V)
A=COS(V)
FUNC1=W-X*A
RETURN
END

DOUBLE PRECISION FUNCTION DR(I)

CONVERT DEGREES TO RADIANS

REAL*8 I
DR=I*3.141592654DO/180DO
RETURN
END

DOUBLE PRECISION FUNCTION RD(I)

CONVERT DEGREES TO RADIANS

REAL*8 I
RD=I*180DO/3.141592654DO
RETURN
END

SUBROUTINE FILER

$INSERT COMMON/MAIN/
$INSERT COMMON/THERM/
REAL*8 CN(9),SPCT(9)
REAL*8 XH,XD
INTEGER DATE
CHARACTER FNAMENAME*35,HEAD*80,NAME*30,DN(10)*2
EQUIVALENCE(ANC,CN(1))
EQUIVALENCE(ANCT,SPCT(1))
DATA DN/'D1 1 , 'D2 1 , 'D3 1 , 'D4 ' , 'D5 1 , 'D6', 'D7', 'D8', 'D9', 'D0'/
TAB=CHAR(137)

C
C...OPEN INPUT FILES
C
C...PARAMETERS AND DAILY SNOWMELT DATA
WRITE(*,*) 'ENTER NAME OF MAIN INPUT FILE: '
READ(*,'(A30)') FNAME
NAME=FNAME
OPEN(9,FILE=FNAME,STATUS='OLD')
C...RAIN CHEMISTRY DATA
WRITE(*,*) 'ENTER NAME OF RAIN INPUT FILE: '
READ(*,'(A30)') FNAME
OPEN(10,FILE=FNAME,STATUS='OLD')
C...ICE LAYER LEVELS DATA
WRITE(*,*) 'ENTER NAME OF ICE ELEV INPUT FILE: '
READ(*,'(A30)') FNAME
OPEN(18,FILE=FNAME,STATUS='OLD')
C...PREVIOUS 10 DAYS OF DISCHARGE DATA FOR MEMORY OF SYSTEM
WRITE(*,*) 'ENTER NAME OF PRE 10 DAYS Q INPUT FILE: '
READ(*,'(A30)') FNAME
OPEN(17,FILE=FNAME,STATUS='OLD')
C...OBSERVED STREAM DATA TO COMPARE IONIC STRENGTHS AND R^2
WRITE(*,*) 'ENTER NAME OF OBS STREAM INPUT FILE: '
READ(*,'(A30)') FNAME

46
OPEN(11,FILE=FNAME,STATUS='OLD')
C...OBSERVED LAKE DATA TO COMPARE R
WRITE(*,*) 'ENTER NAME OF OBS LAKE OUTPUT FILE:'
READ(*, '(A30)') FNAME
OPEN(5,FILE=FNAME,STATUS='OLD')
C...DAILY INFLOW TEMPERATURES, BEGIN W/ J-DAY OF BEGIN OF SIMULATION
WRITE(*,*) 'ENTER NAME OF OBS STREAM TEMP FILE: '
READ(*, '(A30)') FNAME
OPEN(20,FILE=FNAME,STATUS='OLD')
C...THERMOCLINE OBSERVATIONS
WRITE(*,*) 'ENTER NAME OF OBS THERMOCLINE:'
READ(*, '(A30)') FNAME
OPEN(29,FILE=FNAME,STATUS='OLD')
C
C... READ THERMOCLINE ELEVATIONS (MAXIMUM OF 14 VALUES), AND THE J-DAY
IC=1
43 READ(29,*,END=10) XH(IC), XD(IC)
IC=IC+1
GOTO 43
10 CONTINUE
C... OUTPUT FILE NAMES, CREATED BASED ON NAME OF MAIN INPUT FILE
C
C...MAIN OUTPUT FILE (VISUAL)
FNAME='ILWAS>OUT.WY'//'NAME(17:18)///'V'//'NAME(17:30)
OPEN(15,FILE=FNAME,STATUS='UNKNOWN')
C...STREAM CALCULATIONS FILE (TAB DELIMITED FOR MACINTOSH)
FNAME='ILWAS>OUT.WY'//'NAME(17:18)///'S'//'NAME(17:30)
OPEN(13,FILE=FNAME,STATUS='UNKNOWN')
C...STREAM CALC. VS. OBSERVATIONS FILE (TAB DELIMITED FOR MACINTOSH)
FNAME='ILWAS>OUT.WY'//'NAME(17:18)///'RS'//'NAME(17:30)
OPEN(7,FILE=FNAME,STATUS='UNKNOWN')
C...LAKE CALC. VS. OBSERVATIONS FILE (TAB DELIMITED FOR MACINTOSH)
FNAME='ILWAS>OUT.WY'//'NAME(17:18)///'RL'//'NAME(17:30)
OPEN(6,FILE=FNAME,STATUS='UNKNOWN')
C...LAKE CALCULATIONS FILE (TAB DELIMITED FOR MACINTOSH)
FNAME='ILWAS>OUT.WY'//'NAME(17:18)///'L'//'NAME(17:30)
OPEN(12,FILE=FNAME,STATUS='UNKNOWN')
C...IONIC STRENGTH OF STREAM, MACINTOSH FORMAT, CRICKET BAR GRAPH STYLE
FNAME='ILWAS>OUT.WY'//'NAME(17:18)///'I'//'NAME(17:30)
OPEN(14,FILE=FNAME,STATUS='UNKNOWN')
C...LAKE LAYER VOLUMES
FNAME='ILWAS>OUT.WY'//'NAME(17:18)///'VOL'//'NAME(17:30)
OPEN(19,FILE=FNAME,STATUS='UNKNOWN')
C
C... INITIALIZE OUTPUT FILES
C
WRITE(12,1)
1 FORMAT(1H*)
WRITE(12, '(13(A,Al))') 'JDAY', TAB, 'Q', TAB, 'ANC LK', TAB, 'CL LK', TAB,
& 'DOC LK', TAB, 'H LK', TAB, 'NH4 LK', TAB, 'NO3 LK', TAB, 'SBC LK',
& 'TAB', 'SO2 LK', TAB, 'SO4 LK', TAB, 'PH LK',
WRITE(14, '1IONS') TAB, 'J DAY', TAB, 'SBC TAB', 'ANC TAB',
& 'CL TAB', TAB, 'DOC TAB', 'H TAB', 'NO3 TAB', 'SO4 TAB', 'NH4'
WRITE(13,1)
WRITE(13, '(13(A,Al))') 'JDAY', TAB, 'Q TAB', 'ANC ST',
& 'CL ST', TAB, 'DOC ST', TAB, 'H ST'
& TAB, 'NH4 ST', TAB, 'NO3 ST', TAB, 'SBC ST', TAB
& 'SO2 ST', TAB, 'SO4 ST', TAB, '+/- BALANCE ST',
& TAB, 'PH'
C WRITE(7,*) 'JDAY ST', TAB, 'Q ST', TAB, 'ANC ST',
C & 'CL ST', TAB, 'H ST'
C & TAB, 'NH4 ST', TAB, 'NO3 ST', TAB, 'SBC ST', TAB
C 'SI02 ST',TAB,'SO4 ST',TAB,'PH ST'
C 'ANC OB',TAB,'CL OB',TAB,'H OB',TAB,
C 'NH4 OB',TAB,'NO3 OB',TAB,'SBC OB',TAB,'SI02 OB'
C 'SO4 OB',TAB,'PH OB'
C WRITE(6,*)'JDAY LK',TAB,'Q LK',TAB,'ANC LK',
C 'CL LK',TAB,'H LK'
C 'NH4 LK',TAB,'NO3 LK',TAB,'SBC LK',TAB,
C 'SI02 LK',TAB,'SO4 LK',TAB,'PH LK'
C... READ INITIAL DATA VALUES
C... READ OBSERVED VALUES FOR COMPARISONS
C... OBS STREAM DATA TO COMPARE IONIC STRENGTHS, uEQ/L
C 1 = 1
6 READ(11,*,END=11)OSJD(I) ,OSANC(I) ,OSCL(I) ,OSNO3(I) ,
& OSSO4(I) ,OSH(I) ,OSNH4(I) ,OSSBC(I) ,OSIO2(I)
C OSDOC(I)=OSANC(I)+OSCL(I)+OSSO4(I)+OSNO3(I)-OSH(I)-OSNH4(I)
& -OSSBC(I)
1 = 1 + 1
GOTO 6
11 CONTINUE
C... OBS LAKE DATA TO COMPARE R~2, uEQ/L
C 1 = 1
7 READ(5,*,END=12)OLJD(I) ,OLPH(I) ,OLH(I) ,OLNH4(I) ,OLANC(I) ,
& OLCL(I) ,OLNO3(I) ,OLS04(I) ,OLSBC(I) ,OLSIO2(I)
OLDOC(I)=OLANC(I)+OLCL(I)+OLS04(I)+OLNO3(I)-OLH(I)-OLNH4(I)
& -OLSBC(I)
1 = 1 + 1
GOTO 7
12 CONTINUE
C... READ RAIN DATA jday SBC(uEQ/L),CL,NO3,NH4,SO4,H, CALCULATE DOC
C IC=1
READ(10,'(A80)') HEAD
8 READ(10,*,END=9)JDR(IC),SBCR(IC),CLR(IC),NO3R(IC),NH4R(IC)
& ,SO4R(IC),HR(IC),QFALL(IC)
DOCR(IC)=FD(HR(IC),SBCR(IC),NH4R(IC),SO4R(IC),CLR(IC),NO3R(IC))
IC=IC+1
GOTO 8
9 CONTINUE
C... BEGIN TO READ INPUT
C READ(9,'(A80)') HEAD
WRITE(15,'(A80)') HEAD
C. KR, KN NEG. EXPONENTIALS, PO2, PCO2
DO 13 I=1,7
13 READ(9,'(A30)')HEAD
READ(9,*)IMOD
C... AA, BB, PO2 ARE NO LONGER USED
READ(9,*)AA, BB, PO2, PCO2, BAP, JDBIO
C... READ INITIAL COND. OF LAKE, uEQ/L, FIRST ICE, EPILIMNION, THEN HYPOLIMNION
C... DATE, ANC, CL, H, NH4, NO3, SBC, SI02, SO4
READ(9,*)DATI, ANCI, CLI, HI, NH4I, NO3I, SBCI, SI02I, SO4I
READ(9,*)DATE, ANCE, CLE, HE, NH4E, NO3E, SBCE, SI02E, SO4E
READ(9,*) DATE, ANCH, CLH, HH, NH4H, NO3H, SBCH, SiO2H, SO4H
C... NO TEMP CORRECTIONS HERE YET
DOCE=HE+SBCE+NH4E-SO4E-CLE-N03E-ANCE-(10.**(-2.)) /HE
DOCI=HI+SBCI+NH4I-S04I-CLH-N03I-ANCH-(10.**(-2.)) /HI
DOCH=HH+SBCH+NH4H-S04H-CLH-NO3H-ANCH-(10.**(-2.)) /HH
C
C... FIRST DAY, LAST DAY OF SIMULATION, FIRST DAY AND LAST DAY OF MELT
C... AND IF USE MEMORY(=1, USE MEM
C...
READ(9,*) JD1, JDN, JBM, JEM, MEM
C
C... READ ALL DISCHARGES IN METERS CUBED
C... SNOW MELT DATA: jday, Q(M**3)
DO 22 I=JD1, JDN, 1
22 READ(9,*) DAY, QI(I)
C
C... CREATE SNOW HYDROGRAPH SH()
C RAIN STORAGE RS()
C RAIN RUNOFF RR()
C SH = RS SHRS()
C
DO 199 I=1, 365
RS(I)=0.
199 RR(I)=0.
KI=1
DO 200 I=JBM, JEM, 1
IF (I.EQ.JDR(KI)) THEN
RR(I)=QI(I)-QI(I-1)
IF (RR(I).LT.0.0) RR(I)=0.0
IF (RR(I).GT.QFALL(KI)) RR(I)=QFALL(KI)
SH(I)=QI(I)-RR(I)
RS(I)=QFALL(KI)-RR(I)
SHRS(I)=SH(I)-RS(I)
KI=KI+1
ELSE
SH(I)=QI(I)
ENDIF
C WRITE(*, 137) I, TAB, QI(I), TAB, SH(I), TAB, RR(I), TAB, RS(I)
137 FORMAT(I3, A1, 4(F12.2, A1))
200 CONTINUE
C
C... SUM DISCHARGE FOR THE MELT TIME
QTOT=0.0
DO 23 I=JBM, JEM, 1
23 QTOT=QTOT+SH(I)
C
C... BEGIN ADDITION OF DIST. MELT MODIFICATIONS
C
C... READ TOT LOAD OF IONS IN SNOWPACK IN EQUIVALENTS
READ(9,*) SBCT, CLT, NO3T, NH4T, SO4T, HT
C... CALC THE HYDROGEN SUCH THAT THE SNOW PACK MASS HAS CHARGE BALANCE
HT=CLT-NO3T+SO4T-SBCT-NH4T
WRITE(*,*) ' FILER QTOT=', QTOT
DOCT=HT+SBCT+NH4T-SO4T-CLT-NO3T
WRITE(*,*) 'DOC OF SNOWPACK = ', DOCT
C
C... READ # OF SUB UNITS, TOTAL AREA, AND THEIR RESPECTIVE AREAS
READ(9,*) NSU, AREAT, (AREA(IA), IA=1, NSU)
C WRITE(*,*) 'NSU=', NSU
C
C... CREATE NAME OF SUB UNITS FOR THE OUTPUT FILES
IWP=29
DO 111 I=1, NSU
FNAME='ILWAS.OUT.WY'//NAME(17:18)//'>'//DN(I)//NAME(17:30)
OPEN (IWP, FILE=FNAME, STATUS="UNKNOWN")
WRITE (IWP, 1)
WRITE (IWP, '(13(A,1))') 'JDAY', TAB, 'Q', TAB, 'ANC ST', 
   TAB, 'CL ST', TAB, 'DOC ST', TAB, 'H ST'
   ,TAB, 'NH4 ST', TAB, 'NO3 ST', TAB, 'SBC ST', TAB
   , 'SIO2 ST', TAB, 'SO4 ST 1', TAB, '+/- BALANCE ST', 
   TAB, 'PH'
IWP=IWP+1
CONTINUE
C
C... CHECK AREA INPUTS
DO 5 IA=1,NSU
5 AT=AT+AREA(IA)
IF ( (AT/AREAT) .GT.1.1.OR. (AT/AREAT) .LT. .9) THEN
   WRITE (*, *) 'THE AREAS OF THE SUB UNITS DO NOT SUM UP'
   ZZZ=AT/AREAT
   WRITE (*, *) 'RATIO ASUM/AREAT=', ZZZ
   DO 71 I=1,NSU
71 WRITE (*, *) AREA(I)
   WRITE (*, *) 'AREA TOTAL INPUT =', AREAT
   STOP
ENDIF
C
C... READ IN SUB UNITS' SLOPE, ASPECT AND LATITUDE
DO 67 IA=1,NSU
67 READ (9, *) DUMMY, RSLOPE(IA), RASPECT(IA), RLAT(IA)
   , MSEL(IA, 1), MSEL(IA, 2), CF(IA)
CONTINUE
C
C... END ADDITION FOR DIST MELT
C
C... OPEN PREVIOUS 10 DAYS OF DISCHARGE DATA FOR MEMORY OF SYSTEM
C
DO 91 IM=JD1-10,JD1-1
91 QI(IM)=QI(IM)
C
C... READ ICE LEVEL DATA
DO 92 ICE=JD1,JDN
92 READU8, *) I ID, BICE(ICE)
WRITE (*, *) ICE, BICE(ICE)
CONTINUE
C
C... ECHO INPUTS SOFAR
C
C WRITE(15, *) ' ',
C WRITE(15, *) ' KR=10**-', AA
C WRITE(15, *) ' KN=10**-', BB
C WRITE(15, *) ' PCO2=', PCO2
C WRITE(15, *) ' PO2=', PO2
C WRITE(15, *) ' ',
C WRITE(15, *) ' BEGIN DAY=', JD1
C WRITE(15, *) ' END DAY=', JDN
C WRITE(15, *) ' ',
C XXXX=10.5-BICE1
C WRITE(15, *) ' DEPTH OF ICE= ', XXXX
C PH=-LOG10(HE/1000000.)
C WRITE(15, *) ' INITIAL LAKE CONC. uEQ/L, PH=', PH
C WRITE(15, 4)
4 FORMAT (2X, 'ANC', 4X, 'CL', 3X, 'DOC', 4X, 'H', 4X, 
   'NH4', 3X, 'NO3', 3X, 'SBC', 2X, 'SIO2', 3X, 'SO4')
C WRITE(15, 3) ANC, CL, DOC, H, NH4, NO3, SBC, SIO2, SO4, PH
3 FORMAT (14(F5.1,1X),/)
C
RETURN
END
DOUBLE PRECISION FUNCTION FD(H,SBC,NH4,SO4,CL,NO3)
REAL*8 H,SBC,NH4,SO4,CL,NO3
C...CONVERT uEQ/L TO EQ/L (WHICH IS SAME AS MOLES, IN THIS CASE)
H=H/1000000.
SBC=SBC/1000000.
NH4=NH4/1000000.
SO4=SO4/1000000.
CL=CL/1000000.
NO3=NO3/1000000.
C...CALCULATE DOC AND CONVERT BACK TO uEQ/L
FD=(H + SBC + NH4-SO4-CL-N03-(10.**(14.)) /H)*1000000.
H=H*1000000.
SBC=SBC*1000000.
NH4=NH4*1000000.
SO4=SO4*1000000.
CL=CL*1000000.
NO3=NO3*1000000.
RETURN
END

SUBROUTINE RAIN
$INSERT COMMON/MAIN/
REAL*8 CN(9),SPCT(9)
EQUIVALENCE(ANC,CN(1))
EQUIVALENCE(ANCT,SPCT(1))
TAB=CHAR(137)
C... TEST FOR RAIN
C... IF ITS A RAIN DAY, REPLACE SNOWMELT CHEM WITH RAIN CHEM
IFLAG=0
C WRITE(*,*)' JDAY, JDAY OF RAIN',JDAY,JDR(IR)
IF(JDAY.EQ.JDR(IR))THEN
IRAIN1=1
SBC=SBCR(IR)
CL=CLR(IR)
NO3=NO3R(IR)
NH4=NH4R(IR)
SO4=SO4R(IR)
H=HR(IR)
DOC=DOCR(IR)
IF(IYEST.NE.JDAY) THEN
IR=IR+1
ENDIF
IYEST=JDAY
IFLAG=1
WRITE(15,*) 1 RAIN DAY !
ENDIF
WRITE(15,*) 1 JULIAN DAY',JDAY
WRITE(15,*) 1 DISCHARGE M*3',QI(JDAY)
WRITE(15,*)
2 FORMAT(6X,6(F5.1,IX),6X,F5.1,/) 
4 FORMAT(2X,'ANC 1 ,4X,'CL 1 ,3X,'DOC 1 ,4X,'H 1 ,4X,
&'NH4' , 3X, 'NO3' , 3X, 'SBC' , 2X, 'SIO2 1 ,3X, 'SO4')
C... THESE ARE THE INPUT DATA USED TO COMPARE IONIC STRENGTH
IPDOC=DOC
IPSBC=SBC
IPCL=CL
IPNO3=NO3
IPNH4=NH4
IPS04=SO4
IPH=H
RETURN
SUBROUTINE NRWEATH
C
*** NEWTON RAPHSON: WEATHERING ***
C
REMOVED NITROGEN EQUATION; NOW ALL NH4=>H
C
INPUT: PC02, PH, NH4, SBC, CL, SO4, NO3, ANC
C
REEQUILIBRATE: H, NO3, NH4, HCO3, CO3
C
CALCULATE: SI02, SBC
C
INPUT IS IN EQ/L. OUTPUT IS IN EQ/L.
C
EXCEPTION: SI02 IS ALWAYS IN MOLES/L.
C
INTERNAL EQUILIBRIUM CALCULATIONS ARE IN MOLES/L.
C
X, Y, Z = UNKNOWNS

X = H+
Y = NO3-
Z = SI02

Kx = EQUILIBRIUM CONST.

R = ROCK WEATHERING
W = WATER
1, 2 = pK's FOR CARBONATE SYSTEM

PC02 = PRESSURE OF ATM. CO2 GAS

C = NH4 - CL - NO3 - SO4 - DOC

Y1, Y2, Y3 = GOVERNING EQUATIONS = 0.

Y1X = DERIVATIVE OF Y1 W/R/T X
.
.
.
Y3Z = DERIVATIVE OF Y3 W/R/T Z

DV = V(OLD) - M(NEW), etc.

GIVEN: INITIAL ESTIMATES FOR UNKNOWNS

SOLVE BY NEWTON-RAPHSON TO FIND NEW V, W, X, Y, Z

CALCULATE NEW Y1'S AND TEST FOR EQUALITY TO ZERO

IF TEST FAILED, THEN NEW V, W, X, Y, Z VALUES ARE
INITIAL ESTIMATES FOR NEXT ITERATION

SINsert COMMON/MAIN/
REAL*8 CN(9)
REAL*8 HCO3, NO3ST, NO3BI
EQUIVALENCE (ANC, CN(1))
TAB=CHAR(137)
BUG=0
C... OPEN OUTPUT FILE
C  OPEN(16, FILE='IT.LIST', STATUS='UNKNOWN')
C... ITERATION COUNTER
IC=1
C... TOLERANCE OF Y CONVERGENCE TEST
TOL=0.0000000000001

C... RATE CONSTANTS
C... EVALUATE TEMPERATURE
C
C TO = REF. TEMP
C KHO = HENRYS K FOR CO2(g) AT TO
C ST = STREAM TEMP (KELVIN)
C
C KW=10.**(-14.)
C K1=10.**(-6.35)
C K2=10.**(-10.33)
C
K1=10.0**((-3404./ST) - 0.0328*ST + 14.844)
K2=10.0**((-2902./ST) - 0.0238*ST + 6.498)
KH=EXP((-4850/R)*(1/TO-l/ST) + DLOG(KHO))
KW=10.0**(-4470.99/ST - 0.01706*ST + 6.0875)
C
C... THIS IS OLD STUFF, AND ISN'T USED. RUN THE MODEL WITH MEMORY (MEM=1)
IF(MEM.EQ.0)THEN
IF (IMOD.EQ.1)THEN
C... C SERIES USES KR VALUE IN INPUT FILE (EQUILIBRIUM MODEL)
KR=10.**(-AA)
ELSEIF(IMOD.EQ.2)THEN
C... THIS IS WHERE WE LET K=F(Q) (KINETIC MODEL, NO MEM)
CR=0.734
KR=0.30104*QT**(-.63452)
ELSE
C... THIS IS WHERE WE LET K=F(Q) (KINETIC MODEL)
CR=0.721 EXPONENTIAL FIT
KR=3.052*10.**(-3.0)*10.**(4.0773*10.**(5.)*QT)
ENDIF
C
ELSEIF(MEM.EQ.1)THEN
C... HERE WE USE 10 DAY MEMORY IF MEM=1
C IMOD = 1, ACTUAL EQUATION
C IMOD = 2, LOWER BOUND
C IMOD = 3, UPPER BOUND
C BOUNDS ON THE SLOPE OF .002 AND .006
C...NEW STOICHIOMETRY OF 1.2 H : 1.2 SBC, SBC/SIO2 = 1.2
C...NEW KR,89/03/02 ACTUAL
C...NEW KR,89/03/02, UPPER BOUND
C...NEW KR,89/03/02, LOWER BOUND
IF(IMOD.EQ.1)THEN
KR=(0.0081653DO)*10.**(-0.000043561DO * QT)
ELSEIF(IMOD.EQ.2)THEN
KR=(0.015DO)*10.**(-0.000043561DO * QT)
ELSEIF(IMOD.EQ.3)THEN
KR=(0.005DO)*10.**(-0.000043561DO * QT)
ELSEIF(IMOD.EQ.1)THEN
WRITE(*,*)'CHECK INPUT: LOG OR EXP FIT, Q AVERAGE'
STOP
ENDIF
ENDIF
C... NEED DUMMY VALUE FOR SIO2
SIO2=1.0
C... CONVERT UEQ/L TO EQ/L
C
DO 5 JK=2,9
5 CN(JK)=(CN(JK)/1000000.)
C WRITE(*,*) 'ANC=',ANC
C WRITE(*,*) 'CL=',CL
C WRITE(*,*) 'DOC=',DOC
C WRITE(*,*) 'H=',H
C WRITE(*,*) 'NH4=',NH4
C WRITE(*,*) 'NO3=',NO3

53
C... ALL NH4 GOES TO H
C... THEN A % OF NO3 IS RETAINED BY BIOLOGICAL ACTIVITY.
C... THIS VARIABLE IS Biological Activity Percent (BAP)
C...
CAT=NH4+SBC+H
ANI=CL+NO3+SO4
CB=(CAT-ANI)/(CAT+ANI)
C... NITROGEN CYCLE
C... NO3ST = NO3 LEFT IN STREAM
C... NO3BI = NO3 CONSUMED BY BIOTA
IF(JDAY.LT.JDBIO)THEN
NO3 = NO3 + NH4
H = H + 2.* NH4
NH4 = 0.0
ELSEIF(JDAY.GE.JDBIO)THEN
NO3ST = (1.-BAP) * NO3
NO3BI = (BAP) * NO3
NO3 = NO3ST
H = H - NO3BI
H = H + NH4
NH4 = 0.
ENDIF
C=NH4-CL-S04-N03-DOC
C WRITE(*,*)' NRWEATH C= ', C
C... INITIAL CONDITION, UNKNOWNS
X=10.**(-5.1)
Z=SIO2
C... THE Y EQUATIONS
A = K1*KH*PC02
10 Y2= Z*(Z+SBC)**1.2 - KR*X**1.2
IF(BUG.EQ.1)THEN
WRITE(16,*)'INITIAL Y2=',Y2
WRITE(16,*)' Y3=',Y3
WRITE(16,*)' '
ENDIF
C... THE PARTIALS OF Y1,...,Y3 W/R/T X,...,Z
Y2X= -1.2*KR*X**0.2
Y2Z= (Z + SBC)**1.2 + 1.2*Z*(Z+SBC)**0.2
Y3X= l.+A/X**2.+2.*A*K2/X**3.+KW/X**2.
Y3Z= 1.
IF(BUG.EQ.1)THEN
WRITE(16,*)'Y2X=',Y2X
WRITE(16,*)'Y2Z=',Y2Z
WRITE(16,*)'Y3X=',Y3X
WRITE(16,*)'Y3Z=',Y3Z
WRITE(16,*)' '
ENDIF
C... SOLVE FOR DX,DY,DZ
DETA=DET(Y2X,Y2Z,Y3X,Y3Z)
C IF(DETA.LT.1E-10)WRITE(*,*)'DETA=',DETA
DX=DET( Y2,Y2Z,Y3,Y3Z)
DZ=DET(Y2X,Y2 ,Y3X,Y3 )
IF(BUG.EQ.1)THEN
WRITE(16,*)'DETA=',DETA
WRITE(16,*)'DX=',DX
WRITE(16,*)'DZ=',DZ
ENDIF
DX=DX/DETA
DZ=DZ/DETA

IF(BUG.EQ.1) THEN
C    WRITE(16,*)'DX=', DX
C    WRITE(16,*)'DZ=', DZ
C    WRITE(16,*)' ' 
ENDIF
C... SOLVE FOR NEW VALUES OF W,X,Y,Z
IF(BUG.EQ.1) THEN
C    WRITE(16,*)' THE OLD X',X
C    WRITE(16,*)' SOLVING FOR NEW X'
ENDIF
X=BETA(X,DX)
IF(BUG.EQ.1) THEN
C    WRITE(16,*)' THE NEW X',X
C    WRITE(16,*)' ' 
C    WRITE(16,*)' THE OLD Z',Z
C    WRITE(16,*)' SOLVING FOR NEW Z'
ENDIF
C... TEST TO ASSURE NIELHER SIO2 OR SBC CONC. DROPS BELOW ZERO
Z=BETA(Z,DZ)
IF(BUG.EQ.1) THEN
C    WRITE(16,*)' THE NEW Z',Z
ENDIF
C... TEST FOR CONVERGENCE
Y2= Z*(Z+SBC)**1.2 - KR*X**1.2
Y3= Z+ SBC+X+C-A/X-A*K2/X**2.-KW/X

IF(ABS(Y2).LT.TOL.AND.ABS(Y3).LT.TOLJTHEN
    PH=-LOG10(X)
    CO3 = K1*K2*KH*PC02/X**2.
    HCO3 =K1*KH*PC02/X
    H =X
    ANC = HCO3+2*CO3+KW/H-H
    C**  
    ANC= HCO3
    SBC =SBC+Z
    SI02 =Z
C... CONVERT FROM EQUIVALENTS TO MICRO EQUIVALENTS
  DO 6 JK=1,9
    CN(JK)=CN(JK)*1000000. 
  C
C... PROGRAM OUTPUT
  4 FORMAT(2X,'ANC 1 ,4X,'CL',3X,'DOC',4X,'H 1 ,4X,
  &'NH4',3X,'N03 1 ,3X,'SBC',2X,'SI02',3X,' S04')
C... 3 FORMAT(13(F5.1,IX) , /)
  RETURN
ELSE
  36 IC=IC+1
ENDIF
C... RUNAWAY CATCHER
IF(IC.GT.190) THEN
    WRITE(*,*)' MAX ITERATIONS EXCEEDED'
    STOP
ENDIF
GOTO 10
END

DOUBLE PRECISION FUNCTION DET(A,B,C,D)
INTEGER BUG
REAL*8 A,B,C,D
C... CALCULATES THE DETERMINATE OF A 2 X 2 MATRIX
DET=A*D-C*B
RETURN
END
DOUBLE PRECISION FUNCTION BETA(XC,DXC)

$INSERT COMMON/MAIN/
REAL*8 XC,DXC,U
11 U=XC
   IF(BUG.EQ.1) THEN
      WRITE(16,*)'DU=',DXC
   ENDIF
   U=U-DXC
   IF(BUG.EQ.1) THEN
      WRITE(16,*)'U=',U
   ENDIF
   IF(U.LT.0.) THEN
      DXC=DXC/10.
      GOTO 11
   ELSE
      BETA=U
   ENDIF
RETURN
END

SUBROUTINE NROPEN

C
C *** NEWTON RALPHSON: OPEN CARBONATE
C
C UNKNOWNS
C X = H+
C Z = ANC
C
KX = EQUILIBRIUM CONST. H - HENRY'S C02
R = ROCK WEATHERING
W = WATER
1,2 = pK'S FOR CARBONATE SYSTEM
C
PCO2 = PRESSURE OF ATM. C02 GAS
C
C = NH4 -CL -NO3 -SO4 -DOC
C
Y1,Y2 = GOVERNING EQUATIONS = 0.
C
Y1X = DERIVATIVE OF Y1 W/R/T X
C
Y3Z = DERIVATIVE OF Y3 W/R/T Z
C
DV = V(OLD) - M(NEW), etc.
C
GIVEN: INITIAL ESTIMATES FOR UNKNOWNS
C
SOLVE BY NEWTON-RAPHSON METHOD
C
CALCULATE NEW YI'S AND TEST FOR EQUALITY TO ZERO
C
IF TEST FAILED, THEN NEW V,W,X,Y,Z VALUES ARE
C INITIAL ESTIMATES FOR NEXT ITERATION
C
$INSERT COMMON/MAIN/
REAL*8 CN(9)
REAL*8 HCO3
EQUIVALENCE(ANC,CN(1))
TAB=CHAR(137)
BUG=0
C... OPEN OUTPUT FILE
C... ITERATION COUNTER
IC=1
C... TOLERANCE OF Y CONVERGENCE TEST
TOL=0.0000000000001
C... RATE CONSTANTS
C... RATE CONSTANTS
C... EVALUATE TEMPERATURE
C TO = REF. TEMP
C KHO = HENRYS K FOR CO2(g) AT TO
C ST = STREAM TEMP
C...
K1=10.0**((-3404./ST) - 0.0328*ST + 14.844)
K2=10.0**((-2902./ST) - 0.0238*ST + 6.498)
KH=EXP((-4850/R)*(1/TO-1/ST) + DLOG(KHO))
KW=10.0**((-4470.99/ST - 0.01706*ST + 6.0875)
C C... CONVERT uEQ/L TO EQ/L
C DO 5 JK=1,9
5 CN(JK)=(CN(JK) /1000000. )
C C...
C=SBC-NH4-CL-SO4-N03-DOC
C... INITIAL CONDITIONS
X=H
Z=ANC
C... THE Y EQUATIONS
A = K1*KH*PCO2
10 Y2= Z-A/X
Y3= X+C-Z-A*K2/X**2.-KW/X
IF(BUG.EQ.l)THEN
C WRITE(16,*)'INITIAL Y2=',Y2
C WRITE(16,*)' Y3=',Y3
C WRITE(16, *) ' '
ENDIF
C... THE PARTIALS OF Y1,...,Y3 W/R/T X,...,Z
Y2X= A/X**2.
Y2Z= 1.
Y3X= 1.+2.*A*K2/X**3.+KW/X**2.
Y3Z= -1.
IF(BUG.EQ.l)THEN
C WRITE(16,*)'Y2X=',Y2X
C WRITE(16,*)'Y2Z=',Y2Z
C WRITE(16,*)'Y3X=',Y3X
C WRITE(16,*)'Y3Z=',Y3Z
C WRITE(16,*) ' '
ENDIF
C... SOLVE FOR DX,DY,DZ
DETA=DET(Y2X,Y2Z,Y3X,Y3Z)
C IF(DETA.LT.1E-10)WRITE(*,*) 'DETA=',DETA
DX=DET( Y2,Y2Z,Y3,Y3Z)
DZ=DET(Y2X,Y2 ,Y3X,Y3 )
IF(BUG.EQ.l)THEN
C WRITE(16,*) 'DETA=' , DETA
C WRITE(16,*)'DETX=',DX
C WRITE(16,*)'DETZ=',DZ
ENDIF
DX=DX/DETA
DZ=DZ/DETA
IF(BUG.EQ.1)THEN
  WRITE(16,*)'DX=',DX
  WRITE(16,*)'DZ=',DZ
  WRITE(16,*)'
ENDIF
C... SOLVE FOR NEW VALUES OF W,X,Y,Z
IF(BUG.EQ.1)THEN
  WRITE(16,*)' THE OLD X',X
  WRITE(16,*)' SOLVING FOR NEW X'
ENDIF
  X=BETA(X,DX)
IF(BUG.EQ.1)THEN
  WRITE(16,*)' THE NEW X',X
  WRITE(16,*)' THE OLD Z',Z
  WRITE(16,*)' SOLVING FOR NEW Z'
ENDIF
C... TEST TO ASSURE NEITHER SI02 OR SBC CONC. DROPS BELOW ZERO
  Z=BETA(Z,DZ)
IF(BUG.EQ.1)THEN
  WRITE(16,*)'THE NEW Z',Z
ENDIF
C... TEST FOR CONVERGENCE
  Y2= Z-A/X
  Y3= X+C-Z-A*K2/X**2.-KW/X
  IF(ABS(Y2).LT.TOL.AND.ABS(Y3).LT.TOL)THEN
    PH=-LOG10(X)
    CO3 = K1*K2*KH*PC02/X**2.
    HCO3 =Z
    ANC = HCO3+2*CO3+KW/H-H
    C ANC=HCO3
  C... CONVERT FROM EQUIVALENTS TO MICRO EQUIVALENTS
    DO 6 JK=1,9
    6   CN(JK)=CN(JK)*1000000.
    RETURN
ELSE
  WRITE(16,*)' '
  WRITE(16,*)' ITERATION NUMBER ',IC
  WRITE(16,*)' Y2= ',Y2
  WRITE(16,*)' Y3= ',Y3
  WRITE(16,*)' H= ',H
  WRITE(16,*)' X= ',X
  36  IC=IC+1
  C... TEMPORARY STOP
  C STOP
ENDIF
C... RUNAWAY CATCHER
  IF(IC.GT.190)THEN
    WRITE(*,*)' MAX ITERATIONS EXCEEDED OPEN CARB SYS'
    STOP
  ENDIF
GOTO 10
END
SUBROUTINE NRCLOSE
C
C *** NEWTON RALPHSON: CLOSED CARBONATE SYSTEM ***
C
C
C UNKNOWNS
C X = H+
C Z = ANC
C Kx = EQUILIBRIUM CONST. H - HENRY'S CO2
C R - ROCK WEATHERING
C W - WATER
C 1,2 = pK'S FOR CARBONATE SYSTEM
C
C C = NH4 -CL -NO3 -SO4 -DOC
C
C Y1,Y2 = GOVERNING EQUATIONS = 0.
C Y1X = DERIVATIVE OF Y1 W/R/T X
C.
C Y3Z = DERIVATIVE OF Y3 W/R/T Z
C DV = V(OLD) - M(NEW), etc.
C
C GIVEN: INITIAL ESTIMATES FOR UNKNOWNS
C SOLVE BY NEWTON-RAPHSON METHOD
C CALCULATE NEW Y1'S AND TEST FOR EQUALITY TO ZERO
C IF TEST FAILED, THEN NEW V,W,X,Y,Z VALUES ARE
C INITIAL ESTIMATES FOR NEXT ITERATION
C
$INSERT COMMON/MAIN/
REAL*8 CN(9)
REAL*8 HCO3
EQUIVALENCE(ANC,CN(1))
TAB=CHAR(137)
BUG=1
C... OPEN OUTPUT FILE
C OPEN(16,FILE='IT.LIST3',STATUS='UNKNOWN')
C... ITERATION COUNTER
IC=1
C... TOLERANCE OF Y CONVERGENCE TEST
TOL=0.000000000001
C... RATE CONSTANTS
C... EVALUATE TEMPERATURE
C TO = REF. TEMP
C ST = STREAM TEMP
C
C K1=10.0**((-3404./ST) - 0.0328*ST + 14.844)
K2=10.0**((-2902./ST) - 0.0238*ST + 6.498)
KH=EXP((-4850/R)*(1/TO-1/ST) + DLOG(KHO))
KW=10.0**(-4470.99/ST - 0.01706*ST + 6.0875)
C
C... CONVERT UEQ/L TO EQ/L
C DO 5 JK=1,9
5 CN(JK)=(CN(JK)/1000000.)
C
C... CLOSED SYS, CONSERVE TOTAL CARBON (CT)
CT=ANC+ANC*((H/K1+K2/H))
C WRITE(*,*)'CT=',CT
C=SBC-NH4-CL-SO4-NO3-DOC
C... INITIAL CONDITIONS
X=H  
Z=CN(1)  
IF(BUG.EQ.1)THEN  
C WRITE(16,*)'INITIAL X=',X  
C WRITE(16,*)' Z=',Z  
C WRITE(16,*)' '  
ENDIF  
C... THE Y EQUATIONS  
10 Y2= -CT+Z*X/K1+Z*K2*Z/X  
Y3= X+C-KW/X-Z-K2*Z/X  
IF(BUG.EQ.1)THEN  
C WRITE(16,*)'INITIAL Y2=',Y2  
C WRITE(16,*)' Y3=',Y3  
C WRITE(16,*)' '  
ENDIF  
C... THE PARTIALS OF Y1,...,Y3 W/R/T X,...,Z  
Y2X= Z/K1+K2*Z/X**2.  
Y2Z= X/K1+1.+K2/X  
Y3X= 1.+KW/X**2.+K2*Z/X**2.  
Y3Z= -1.-K2/X  
C IF(BUG.EQ.1)THEN  
C WRITE(16,*)'Y2X=',Y2X  
C WRITE(16,*)'Y2Z=',Y2Z  
C WRITE(16,*)'Y3X=',Y3X  
C WRITE(16,*)'Y3Z=',Y3Z  
C WRITE(16,*)' '  
C ENDIF  
C... SOLVE FOR DX,DY,DZ  
DETA=DET(Y2X,Y2Z,Y3X,Y3Z)  
IF(ABS(DETA).LT.1E-10)WRITE('*,*)'DETA=',DETA  
DX=DET( Y2,Y2Z,Y3,Y3Z)  
DZ=DET (Y2X,Y2 ,Y3X,Y3 )  
IF(BUG.EQ.1)THEN  
C WRITE(16,*)'DETA=',DETA  
C WRITE(16,*)'DETX=',DX  
C WRITE(16,*)'DETZ=',DZ  
ENDIF  
DX=DX/DETA  
DZ=DZ/DETA  
C IF(BUG.EQ.1)THEN  
C WRITE(16,*)'THE NEW X',X  
C WRITE(16,*)' '  
C ENDIF  
C... TEST TO ASSURE NIETHER SIO? OR SBC CONC. DROPS BELOW ZERO  
Z=BETA(Z,DZ)  
C IF(BUG.EQ.1)THEN  
C WRITE(16,*)'THE NEW Z',Z  
C ENDIF  
C... TEST FOR CONVERGENCE  
Y2= -CT+Z*X/K1+Z*K2*Z/X  
Y3= X+C-KW/X-Z-K2*Z/X  
60
IF (BUG.EQ.1) THEN
  WRITE(16,*)' NEW Y2=', Y2
  WRITE(16,*)' Y3=', Y3
ENDIF

IF (ABS(Y2).LT.TOL.AND.ABS(Y3).LT.TOL) THEN
  WRITE(*,*)' FINAL Y2=', Y2
  WRITE(*,*)' Y3=', Y3
  WRITE(*,*)' X=', X
  WRITE(*,*)' Z=', Z
  HC03 = Z
  CO3 = K2*HC03
  H = X
  ANC = HC03 + 2*CO3 + KW/H - H
  WRITE(16,*) ' HC03 = ', HC03
  WRITE(16,*) ' CO3 = ', CO3
  WRITE(16,*) ' H = ', H
  WRITE(16,*) ' ANC = ', ANC
  PH = -LOG10(X)
  C... CONVERT FROM EQUIVALENTS TO MICRO EQUIVALENTS
  DO 6 JK=1,9
    CN(JK) = CN(JK)*1000000.
  RETURN
ELS
  WRITE(16,*) ' ITERATION NUMBER ', IC
  WRITE(16,*) ' Y2= ', Y2
  WRITE(16,*) ' Y3= ', Y3
  IC = IC + 1
C... TEMPORARY STOP
  STOP
ENDIF
C... RUNAWAY CATCHER
IF (IC.GT.1490) THEN
  WRITE(*,*)' MAX ITERATIONS EXCEEDED CLOSED CARB SYS'
  STOP
ENDIF
GOTO 10
END

SUBROUTINE REQ(VICE)
C... REEQUILIBRATE LAKE EPI LAYER
C... NROPEN = OPEN SYSTEM
C... NRCLOSE = CLOSED SYSTEM
C
  IF (VICE.LE.0.0) THEN
    CALL NROPEN
  ELSE
    CALL NRCLOSE
  ENDIF
  RETURN
END

SUBROUTINE LAKE(IKI)
C TAKES PREVIOUS CONC OF EP. AND HY. LAYERS AND MIXES THEM
C AS THE THERMOCLINE CHANGES.
C VT= VOL TOTAL
C V2= VOL EPILIMNION, NEW (TODAY)
C V2= VOL EPILIMNION, OLD (YESTERDAY)
C VH= VOL HYPO NEW
C VHI= VOL HYPO OLD
C VHD= DIFFERENCE IN HYPO VOLUME DUE TO CHANGE IN THERMOCLINE

   }
DIV = DIFFERENCE IN ICE VOLUME DUE TO MELT/FREEZING
VICE = ICE VOLUME
BIce() = ELEVATION OF ICE BOUNDARY ABOVE LAKE BOTTOM
CI1() = CHEM CONC. IN ICE
CI2() = NEW DAY CONC. IN ICE

TAKES CONC. VALUES IN COMMON AND MIXES THEM WITH CONC.
OF LAKE FROM PREVIOUS TIME STEP, IN ARRAY CE1,CH1, WHICH IS INITIALIZED
IN THE FIRST CALL TO THIS SUB FROM MAIN, WHEN ARGUMENT I = 1.

ARRAY CN IS EQUIVALENCED WITH THE IN COMING CONCENTRATIONS
AND THE OUT GOING NEW CONCENTRATIONS.

ARRAY CE1,CH1 IS THE MEMORY OF THE LAKE. HERE WE STORE FOR ONE TIME
STEP THE LAST VALUES OF CONCENTRATION OF THE LAKE. WHEN IKI = 1
THE CAT/ANION VALUES ARE READ INTO CE1,CH1

SUBROUTINE THERM WILL DETERMINE THE HEIGHT OF THE THERMOCLINE
BASED ON THE DATE, AND RETURN VOLUME OF THE EPILIMNION.

$INSERT COMMON/MAIN/
REAL*8 CN(9)
EQUIVALENCE(ANC,CN(1))
TAB=CHAR(137)
VT=179600.0
C
C... IF BLOCK EXECUTED ONCE IN BEGINNING OF SIMULATION, TO INITIALIZE
C... LAKE CONCENTRATIONS, AND OPEN ANY FILES
C
IF(IKI.EQ.1)THEN
OPEN(29,FILE='ILWAS>OUTPUT>MIX.ANC',STATUS='UNKNOWN')
C... FOLLOWING OPEN STATEMENTS USED TO CHECK MIXING OF LAKE.
WRITE(29,*)(JDAY,TAB,'EPI VOL',TAB,'E CON ',TAB,'EPI VOLAI'
&TAB,'EC AI',TAB,'EV AH',TAB,'EC AH',TAB,'STR VOL',TAB,
&TAB,'STR CONC',TAB,'SV AM',TAB,'SC AM'
OPEN(30,FILE='ILWAS>OUTPUT>MIX.SIO2',STATUS='UNKNOWN')
OPEN(31,FILE='ILWAS>OUTPUT>MIX.SBC',STATUS='UNKNOWN')
OPEN(32,FILE='ILWAS>OUTPUT>MIX.H',STATUS='UNKNOWN')
OPEN(33,FILE='ILWAS>OUTPUT>MIX.CL',STATUS='UNKNOWN')
OPEN(34,FILE='ILWAS>OUTPUT>MIX.NO3',STATUS='UNKNOWN')
OPEN(35,FILE='ILWAS>OUTPUT>MIX.NH4',STATUS='UNKNOWN')
OPEN(36,FILE='ILWAS>OUTPUT>MIX.SO4',STATUS='UNKNOWN')
C... ICE
CI1(1)=ANC1
CI1(2)=CLI
CI1(3)=DOCI
CI1(4)=HI
CI1(5)=NH4I
CI1(6)=NO3I
CI1(7)=SBCI
CI1(8)=SI02I
CI1(9)=SO4I
C... EPILIMNION
CE1(1)=ANCE
CE1(2)=CLE
CE1(3)=DOCE
CE1(4)=HE
CE1(5)=NH4E
CE1(6)=NO3E
CE1(7)=SBCE
CE1(8)=SI02E
CE1(9)=SO4E
C... HYPOLIMNION
CH1(1)=ANCH
CH1(2)=CLH
CH1(3)=DOCH
CH1(4)=HH
CH1(5)=NH4H
CH1(6)=NO3H
CH1(7)=SBCH
CH1(8)=SI02H
CH1(9)=SO4H
RETURN
ENDIF
C
C... CALL THERMOCLINE TO FIND VOLUME OF EPILIMNION AND ICE LAYER
C
CALL THERM(JDAY,BICE(JDAY),VE,VICE)
VH=VT-VE-VICE
IF(VH.LT.0.0)VH=0.
C
C... CHECK THE MIXING MASS BALANCE HERE
C
IA=1
WRITE(29,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
IA=2
WRITE(30,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
IA=3
WRITE(31,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
IA=4
WRITE(32,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
IA=5
WRITE(33,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
IA=6
WRITE(34,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
IA=7
WRITE(35,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
IA=8
WRITE(36,20)JDAY,TAB,CI1(IA),TAB,VICE,TAB,CE1(IA),TAB,VE,TAB,
&CH1(IA),TAB,VH,TAB,CN(IA),TAB,QI(JDAY)
C
C... SET INITIAL VALUES FOR VOLUMES, SO NO ICE MELT OR THERMOCLINE MOVEMENT
C... OCCUR ON FIRST DAY
C... VICE1 IS YESTERDAY'S ICE VOLUME
C... VH1 IS YESTERDAYS HYPOLIMNION VOLUME
C
IF(JDAY.EQ.JD1)THEN
VICE1=VICE
VE1=VE
VH1=VH
ENDIF
C
C... THE FOLLOWING CHANGES IN VOLUMES CONTROL THE LAKE LAYER MIXING
C
VHD=VH1-VH
DIV=VICE-VICE1
CTW
EVBM=VE1
ECBM=CE1(1)
C WRITE(*,*)'VE1 BM=',VE1
CTW
C.
C... BEGIN ICE LAYER/EPILIMNION INTERFACE MIXING
C.
IF(DIV.LT.-0.1)THEN
C. ADD TO EP, SUBTRACT FROM ICE
DO 8 I=1,9
  CE2(I)=(CE1(I)*VE1+CI1(I)*ABS(DIV))/(VE1+ABS(DIV))
  CE1(I)=CE2(I)
  IF((VICE1-ABS(DIV)).LE.0.)THEN
    CI1(I)=0.0
  ELSE
    CI2(I)=(CI1(I)*VICE1-CI1(I)*ABS(DIV))/(VICE1-ABS(DIV))
    CI1(I)=CI2(I)
  ENDIF
8 CONTINUE
VE1=VE1+ABS(DIV)
ELSEIF(DIV.GT.0.1) THEN
C. ADD TO ICE, SUBTRACT FROM EP
DO 9 I=1,9
  CI2(I)=(CI1(I)*VICE1+CE1(I)*ABS(DIV))/(VICE1+DIV)
  CI1(I)=CI2(I)
  IF((VE1-DIV).LT.0.0)THEN
    CE1(I)=0.0
  ELSE
    CE2(I)=(CE1(I)*VE1-CE1(I)*ABS(DIV))/(VE1-DIV)
    CE1(I)=CE2(I)
  ENDIF
9 CONTINUE
VE1=VE1-ABS(DIV)
C. NEXT LINE TO CORRECT POSSIBLE ROUND OFF
IF(VE1.LT.0.0)VE1=0.0
ENDIF
C. REPLACE OLD VALUES WITH NEW VALUES, ALSO NO CHANGE IN ICE LAYER
VICE1=VICE
CTW
EVAI=VE1
ECAI=CE1(1)
C WRITE(*,*)'VE1 AI=',VE1
CTW
C.
C. . . END ICE
C.
C... BEGIN EPI/HYPO INTERFACE MIXING
C.
C WRITE(*,*)'VHD= ',VHD
IF(VHD.GT.0.1)THEN
C. ADD TO EP, SUBTRACT FROM HY
DO 10 I=1,9
  CE2(I)=(CE1(I)*VE1+CH1(I)*VHD)/(VE1+VHD)
  CE1(I)=CE2(I)
  IF((VH1-VHD).LE.0.)THEN
    CH1(I)=0.0
  ELSE
    CH2(I)=(CH1(I)*VH1-CH1(I)*VHD)/(VH1-VHD)
    CH1(I)=CH2(I)
  ENDIF
10 CONTINUE
64
VE1 = VE1 + VHD
ELSEIF (VHD.LT.-0.1) THEN
C. ADD TO HY, SUBTRACT FROM EP
DO 11 I=1,9
  CH2(I) = (CH1(I)*VH1 + CE1(I)*ABS(VHD)) / (VH1 + ABS(VHD))
  CH1(I) = CH2(I)
  IF((VE1-ABS(VHD)).LT.0.0) THEN
    CE1(I) = 0.0
    C... ABOVE CASE SHOULD NEVER HAPPEN, THERMOCLINE AND ICE/EPI INTERFACE MEET
    ELSE
      CE2(I) = (CE1(I)*VE1 - CE1(I)*ABS(VHD)) / (VE1 - ABS(VHD))
      CE1(I) = CE2(I)
    ENDIF
  CONTINUE
VE1 = VE1 + VHD
ENDIF
C. REPLACE OLD VALUES WITH NEW VALUES, ALSO NO CHANGE IN THERMOCLINE CASE
VE1 = VE
VH1 = VH
C... CHECK VOLUMES
WRITE(19,14) JDAY, TAB, VT, TAB, VICE, TAB, VE1, TAB, VH1
C... END CHECK VOLUMES
CTW
EVBM = VE1
ECAH = CE1(1)
C... END EPI.HYPO INTERFACE MIXING
C... DO THE MIXING OF THE INFLOW
C... CTW
SVBM = QI(JDAY)
SCBM = CN(1)
CTW
VT = VE + QI(JDAY)
DO 100 I=1,9
  CE1(I) = (QI(JDAY) * CN(I) + VE1 * CE1(I)) / VT
  CN(I) = CE1(I)
100
C... RE-EQUILIBRATE LAKE EPI LAYER
CALL REQ(VICE)
WRITE(29,20) JDAY, TAB, CN(1), TAB, CN(2), TAB, CN(3), TAB, CN(4)
C 6, TAB, CN(5), TAB, CN(6), TAB, CN(7), TAB, CN(8), TAB, CN(9)
C... RE-EQUILIBRATE LAKE EPI LAYER
CALL REQ(VICE)
WRITE(29,20) JDAY, TAB, CN(1), TAB, CN(2), TAB, CN(3), TAB, CN(4)
C 6, TAB, CN(5), TAB, CN(6), TAB, CN(7), TAB, CN(8), TAB, CN(9)
C... CTW
SVAM = VT
SCAM = CN(1)
C
WRITE(29,20) JDAY, TAB, EVBM, TAB, ECAH
C 6, TAB, EVAI, TAB, ECAI
C 6, TAB, EVBM, TAB, SCBM
C 6, TAB, SVBM, TAB, SCAM
CTW
C
14 FORMAT (I3, A1, F10.0, A1, F10.0, A1, F10.0, A1, F15.0, A1, F15.0)
15 FORMAT (I3, 3X, 5(4X, F10.1))
IF (JDAY.EQ.OLJD(ILOD)) THEN
    OLH(ILOD) = OLH(ILOD)/1000000.
    PH = -LOG10(H/1000000.)
    OLH(ILOD) = OLH(ILOD)*1000000.
WRITE(6,43) JDAY, TAB, QI(JDAY), TAB, ANC, TAB, CL, TAB, H
& TAB, NH, TAB, NO3, TAB, SBC, TAB, SIO2, TAB, SO4, TAB, PH
& TAB, OLANC(ILOD), TAB, OLCL(ILOD), TAB, OLH(ILOD), TAB,
& OLNH4(ILOD), TAB, OLN03(ILOD), TAB, OLSCB(ILOD), TAB, OLSBC(ILOD),
& TAB, OLS04(ILOD), TAB, OLPH(ILOD)
ILOD = ILOD + 1
ENDIF
C WRITE(15,*) ' EPILIMNION VOLUME =', VE
C WRITE(15,*) ' EPILIMNION CONC. AFTER MIXING, uEQ/L'
C WRITE(15,4)
C WRITE(15,42) ANC, CL, DOC, H,
& NH4, NO3, SBC, SIO2, SO4
XPH = -LOG10(H/1000000)
WRITE(12,41) JDAY, TAB, QI(JDAY), TAB, ANC, TAB, CL, TAB, DOC, TAB, H, TAB,
& NH4, TAB, NO3, TAB, SBC, TAB, SIO2, TAB, SO4, TAB, XPH
41 FORMAT(I3, A1, F10.1, A1, F10.1, A1, F10.1, A1, F10.1, A1, F10.1,
& A1, F10.1, A1, F10.1, A1, F10.1, A1, F10.1, A1, F10.1,
& A1, F10.1, A1, F10.1, A1, F10.1, A1, F10.1, A1, F10.1)
42 FORMAT(F10.1, F10.1, F10.1, F10.1, F10.1,
& F10.1, F10.1, F10.1, F10.1, F10.1)
& 'NH4', 3X, 'NO3', 3X, 'SBC', 2X, 'SIO2', 3X, 'SO4')
3 FORMAT(9(F5.1,IX),'/')
RETURN
END

SUBROUTINE THERM(JDAY, AL, VL, VICE)
C ... AL = HEIGHT OF BOTTOM OF ICE FROM LAKE BOTTOM
C ... INSERT COMMON/THERM/
C XH IS OBS THERMOCLINE HEIGHT ON JULIAN DAY XD
C REAL*8 VL, AL, VICE, XH, XD
REAL TC
C GIVEN DAY, RETURN VOLUME OF EPILIMNION(VL) AND ICE(VICE)
C DO 10, IC=1,14
   IF(JDAY.GT.XD(IC)) I=IC
10 CONTINUE
   TC = TCE(I, JDAY)
C ... TC IS ELEVATION OF THERMOCLINE
C ... AL IS ELEVATION OF BOTTOM OF ICE LAYER
C ... AL CANNOT BE LOWER THAN TC
   IF(AL.LT.TC) THEN
      WRITE(*,*) ' ERROR IN ICE AND THERMOCLINE ELEVATIONS !!'
      STOP
   ENDIF

66
IF (TC.LT..5) I=1
IF (TC.GE..5) I=2
IF (TC.GE.1.5) I=3
IF (TC.GE.2.5) I=4
IF (TC.GE.3.5) I=5
IF (TC.GE.4.5) I=6
IF (TC.GE.5.5) I=7
IF (TC.GE.6.5) I=8
IF (TC.GE.7.5) I=9
IF (TC.GE.8.5) I=10
IF (TC.GE.9.5) I=11
VL=VOL(I,TC)
C WRITE(*,*) ' TC ELEV = ', TC
C WRITE(*,*) ' EPI VOL = ', VL
TC=AL
IF (TC.LT..5) I=1
IF (TC.GE..5) I=2
IF (TC.GE.1.5) I=3
IF (TC.GE.2.5) I=4
IF (TC.GE.3.5) I=5
IF (TC.GE.4.5) I=6
IF (TC.GE.5.5) I=7
IF (TC.GE.6.5) I=8
IF (TC.GE.7.5) I=9
IF (TC.GE.8.5) I=10
IF (TC.GE.9.5) I=11
VICE=VOL(I,TC)
C WRITE(*,*) ' ICE ELV = ', TC
C WRITE(*,*) ' ICE VOL = ', VICE
VL=VL-VICE
C WRITE(*,*) ' NEW EPI VOL = ', VL
RETURN
END
C...
FUNCTION TCE(I,JDAY)
$INSERT COMMON/THERM/
REAL*8 XH,XD
C... INTERPOLATE THERMOCLINE ELEVATION BASED ON INPUT VALUES OF
C... THERMOCLINE HEIGHT AND CORRESPONDING DAYS
XM=(XH(I+1)-XH(I))/(XD(I+1)-XD(I))
TCE=XH(I+1)-XM*(XD(I+1)-FLOAT(JDAY))
RETURN
END
C...
FUNCTION VOL(I,TC)
DIMENSION T(12),V(12)
DATA T/0.,.5,1.5,,2.5,3.5,4.5,5.5,6.5,7.5,8.5,9.5,10.5/,
& 13000.,15160.,17960./
C... INTERPOLATE VOLUME OF THE HYPOLIMNION FROM THE TC ELEVATION
M=(V(I+1)-V(I))/(T(I+1)-T(I))
VOL=V(I+1)-M*(T(I+1)-TC)
C... EPILIMNION VOL = TOTAL VOL - HYPOLIMNION VOL
VOL=179600-VOL
C WRITE(*,*) ' VOL',VOL
RETURN
END
SUBROUTINE IONIC
$INSERT COMMON/MAIN/
67
C... WRITES IN A SPECIFIED FORMAT FOR MAKING BAR GRAPHS IN CRICKET GRAPH
C... A MACINTOSH APPLICATION. USEFUL FOR COMPARING IONIC STRENGTHS
C
C Z = 0.
C.. IF (IFLAG.EQ.1) THEN
WRITE (14,1) TAB, JDAY, TAB, IPSBC, TAB, Z, TAB
& Z, TAB, Z, TAB, IPH, TAB, Z, TAB, Z, TAB, IPNH4
1 FORMAT (16HCATIONS INPUT(R), A1, I3, 8(A1, F14.2))
WRITE (14, 3) TAB, JDAY, TAB, Z, TAB, Z, TAB
& IPCL, TAB, IPDOC, TAB, Z, TAB, IPNO3, TAB, IPSO4, TAB, Z
3 FORMAT (16H ANIONS INPUT(R), A1, I3, 8(A1, F14.2))
ELSE
WRITE (14, 2) TAB, JDAY, TAB, IPSBC, TAB, Z, TAB
& Z, TAB, Z, TAB, IPH, TAB, Z, TAB, Z, TAB, IPNH4
2 FORMAT (16HCATIONS INPUT(S), A1, I3, 8(A1, F14.2))
WRITE (14, 4) TAB, JDAY, TAB, Z, TAB, Z, TAB
& IPCL, TAB, IPDOC, TAB, Z, TAB, IPNO3, TAB, IPSO4, TAB, Z
4 FORMAT (16H ANIONS INPUT(S), A1, I3, 8(A1, F14.2))
ENDIF
C.. WRITE(*,*)'JDAY 1, JDAY, ' OSJD(',, IO, ') ', OSJD(IO)
IF (JDAY.EQ.OSJD(IO)) THEN
WRITE (14, 7) TAB, OSJD(IO), TAB, OSSBC(IO), TAB, Z, TAB
& Z, TAB, Z, TAB, OSH(IO), TAB, Z, TAB, Z, TAB, OSNH4(IO)
7 FORMAT (11HCATIONS OBS, A1, I3, 8(A1, F14.2))
WRITE (14, 8) TAB, OSJD(IO), TAB, Z, TAB, OSANC(IO), TAB
& OSCL(IO), TAB, OSDOC(IO), TAB, Z, TAB, OSNO3(IO), TAB
& OSSO4(IO), TAB, Z
IO=IO+1
ENDIF
WRITE (14,*) TAB
8 FORMAT (11H ANIONS OBS, A1, I3, 8(A1, F14.2))
RETURN
END