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**An Inverse Approach to the Construction of Fracture  
Hydrology Models Conditioned by Geophysical Data—  
An Example from the Validation Exercises at the Stripa Mine**

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## ABSTRACT

One approach for the construction of fracture flow models is to collect statistical data about the geometry and hydraulic apertures of the fractures and use this data to construct statistically identical realizations of the fracture network for fluid flow analysis. We have found that this approach has two major problems. One is that an extremely small percentage of visible fractures may be hydrologically active. The other is that, on any scale you are interested in characterizing usually a small number of large features dominate the behavior (Long *et al.*, 1989). To overcome these problems we are proposing an approach in which the model is strongly conditioned by geology and geophysics. Tomography is used to identify the large features. The hydraulic behavior of these features is then obtained using an inverse technique called "simulated annealing." The first application of this approach has been at the Stripa Mine in Sweden as part of the Stripa Project. Within this effort, we built a model to predict the inflow to the Simulated Drift Experiment (SDE), i.e. inflow to six parallel, closely spaced holes (two-holes), using a preliminary set of data collected in five other holes, the N- and W-holes. We predict a mean total flow of approximately 3.1 (*l/min*) into the six holes (two-holes) with a coefficient of variation nearly unity and a prediction error of about 4.6 *l/min*. The actual measured inflow is close to 2 *l/min*.

## 1.0. INTRODUCTION

The fundamental problem in modeling fracture hydrology is that the medium does not behave as a continuum. Parts of the rock have no hydraulic communication with other parts. In the case where the matrix rock can be considered impermeable, this fundamental behavior is governed by the geometry of the fracture network. If a connected cluster of conductive fractures links two points, there can be flow between the two points. If not, then there is no flow. This heterogeneous connectivity is very commonly observed in fractured rock. As a consequence of this observation, attempts at building a fracture hydrology model should be focussed on reproducing, this fundamental behavior. Several attempts have been made to find appropriate fracture networks by statistically reproducing the geometry of the fracture network (Long and Billaux, 1987; Khaleel, 1989; Billaux *et al.*, 1989; La Pointe and Hudson, 1985; among others). This involves determining a statistical rule for locating fractures, determining their orientation, size and conductivity. Then the interconnections between the fractures can be found and flow patterns calculated.

For example, an interpretation was made of the data at the Fanay-Augères mine in France (Billaux *et al.*, 1989) using data which included fracture trace maps from drift walls giving fracture location, orientation, trace length, coatings, roughness etc.; fracture logs from boreholes giving fracture location, orientation, coatings, roughness etc.; and single-hole packer tests. The analysis was based on assuming the fractures were disc-shaped and uniformly permeable in their plane. In order to represent the clustering of fractures, the arrangement of fractures in space was a complex Poisson process called the Parent-Daughter process. Geostatistical techniques were used to fit the parameters of the model based on data on trace lengths, orientation and fracture frequency. A pattern of fractures in a 100 m cube was determined.

Conclusions from this effort were very striking. First we found that the use of one- and two-dimensional data to infer three-dimensional geometry is a highly non-unique problem. Many three-dimensional geometries can account for the same one- and two-dimensional data. Second,

no matter how the three-dimensional geometry was determined, there were far too many fractures to account for the lack of connectivity known to exist. If all the fractures were present and hydrologically active, the medium would have behaved like an equivalent continuum. However, cross-hole hydrologic and tracer test results showed that this was definitely not the case. It must be the case at Fanay-Augères that only a very, very small part of the fracture network conducts fluid. We suspect that many cases are similar to Fanay-Augères in that the fractures are relatively pervasive and the real problem is finding what parts of the system actually conduct.

Fanay-Augères offered one other key fact in this regard. The two drifts were mapped in this mine: one wet (S1), one dry (S2). For both drifts the fracture geometry seemed to indicate highly connected fracture networks. However, in the wet drift, a major fault ran through the block of rock surrounding this portion of the drift. It seems that the hydrology of the site is controlled by major features i.e. fracture zones. This observation is certainly not confined to Fanay-Augères. Similar conclusions can be drawn from Olsson *et al.* (1990) (Carlsten *et al.*, 1988) at Stripa and Hsieh *et al.* (1985) who performed cross-hole hydraulic tests in a ubiquitously fractured granite but found that certain zones had no hydraulic connection to others. It is possible to find many other references which describe the same phenomena. From these facts we infer that the hydrology of many fractured sites is controlled by a finite number of major features.

Our experience so far has indicated that focussing on the details of fracture geometry statistics is equivalent to "not seeing the forest for the trees." If fracture zones control the hydrology, then efforts should first be aimed directly at locating and characterizing fracture zones. In this method we place emphasis on determining the hydrologic characteristics of the zones conditioned by the geometry of the fractures rather than letting predicted geometry predict hydrology. We expect that zones are not continuous and that the permeability structure within the zones is complex. Therefore, we propose a "top - down" approach for building a fracture hydrology model which

1. locates important zones

2. characterizes these zones
3. characterizes the "background" fractures.

To obtain a model of the first order hydrologic behavior, we start by modeling the major hydrologic features which, in the case of Stripa, are assumed to be fracture zones. We assume these zones are heterogeneous and we then use a hydrologic inversion to find the pattern of conductance within the zones. Thus in this case we are including two levels of detail: the zones themselves and the conductance pattern within the zone. The zones are considered to be very important and the actual location of the zones is considered carefully. The pattern of conductance within the zones is considered second order. These are represented by an equivalent lattice of conductors which lie on a grid called a "template."

For location of the zones, we rely heavily on the recent advances in geophysics that have allowed fractures zones to be "seen" inside the rock. Thus the model is conditioned by geophysical information. Then, based on these we build a hydrologic conceptual model of the the rock which we call a template. The template should contain all the likely major conductors and is the basis for fluid flow calculation. We infer as much as possible about the qualitative hydrologic attributes of the zones through geology and geomechanics. Finally, we perform a hydrologic inversion on the model. We throw away conductors in a manner that conditions the model to observe well test behavior. In other words, within the template we identify patterns of conductance that can explain the observed hydraulic behavior. We then use simulated annealing to arrange the conductances such that they explain observed distributions of head, observed fluxes, or observed tracer test results.

### **1.1. Background**

Investigations related to the geologic storage of nuclear waste have been ongoing at the Stripa Mine in Sweden for more than ten years. The latest of these investigations is called Phase III and is sponsored by OECD Nuclear Energy Association (NEA) as an international cooperative effort managed by the Swedish Nuclear Fuel and Waste Management Company (SKB).

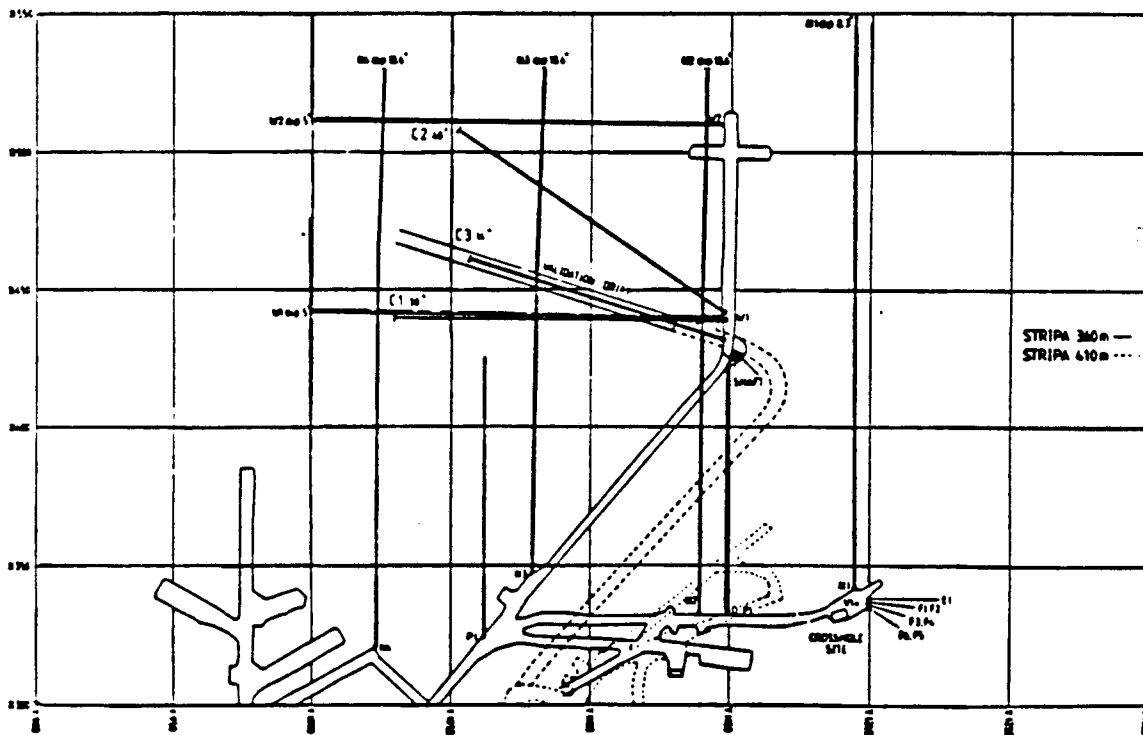


The Stripa Phase III project includes the Site Characterization and Validation (SCV) experiment, which is designed to test current abilities to characterize fractured rock before it is used for nuclear waste storage. The effort is centered on a block of rock 150 m long  $\times$  100 m wide  $\times$  50 m deep in size at a depth of about 330 m. The block lies between previous experimental sites, the Macroporosity/Buffer Mass Test site and the 3-D Migration site (Figure 1.1). The SCV work is divided into two cycles of three stages each: data-gathering, prediction, and validation. The first stage of work has included drilling of 6 boreholes (N2, N3, N4, W1, W2, and V3) and taking measurements of geology, fracture characteristics, stress, single borehole geophysical logging, radar, seismics, and hydrogeology (Olsson *et al.*, 1990). Olsson *et al.* gives background information describing the site and the results of the first cycle of data-gathering and prediction. The analysis presented here are based almost entirely on the data documented in Olsson *et al.* (1990).

## 1.2. Goals

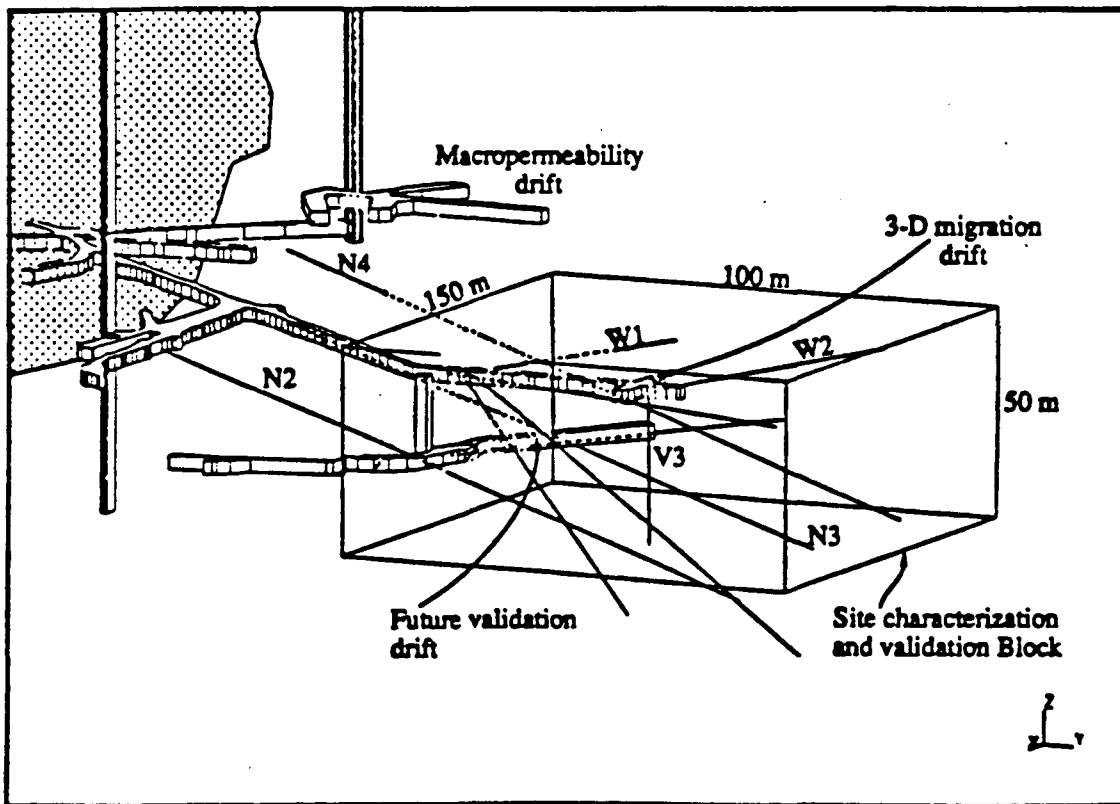
Hydrologic prediction of groundwater flow and transport is part of the work scope for the Stripa Phase III Project (SKB, 1987). Current plans call for two prediction cycles. The first cycle is focused on the prediction of inflow into a clustered group of boreholes (the D-holes) which are drilled on the periphery of the future "Validation Drift." The D-holes are drilled such that five of them lie on the periphery of the future Validation Drift and one lies in the center (Figure 1.2). The measurement of flow into the D-holes is called the Simulated Drift Experiment. In the second cycle, the Validation Drift is excavated and then the inflow into the drift is predicted. In both cases, actual measurements of inflow will be compared to the predictions.

This paper covers LBL's preliminary prediction of flow into the D-holes. The work should be read as an example of an approach to the problem. The actual results are severely impacted by the fact that controlled cross-hole hydrologic tests were unavailable. For the purpose of demonstration, we have inferred from available data what a cross-hole test might show and used this synthetic data to make a preliminary calculation of the inflow into the D-holes.



XBL 903-847

Figure 1.1. Plan view of the SCV site showing the location of the C-holes and the Validation Drift.



XBL 903-849

Figure 1.2. Perspective view of the SCV block. Dotted area in the upper left is the mined out stopes (after J. Gale).

### 1.3. Validation

Testing our predictions of inflow into D-holes against the measured values is part of an exercise aimed at validating fracture flow codes. The first component is the calculation of prediction error. That is, we use the model to make a series of predictions. For each prediction we obtain a prediction error by comparing the calculated result to the measured result. The root mean square of these errors is called the prediction error. Thus, we validate the model by estimating how well the model makes predictions. For example, we could use the model to predict the inflow into ten different boreholes for which the inflow had been measured. By comparing the measurements of inflow to the predicted values, we can calculate a prediction error. Now, we can use the model to predict the flow into an eleventh hole for which the inflow is not known. The prediction error then provides an estimate of how good this prediction is, i.e. how valid the model is.

This approach is straight forward. The more numerous and diverse types of predictions that can be included in the estimate of prediction error, the more stringently "validated" the model can be. Although the model may work well to predict flow under one type of boundary condition, it may not predict well for a different type of boundary conditions. However, if it predicts well for two sets, then it is more likely to predict well under the third, even better for three, etc., etc.

A limitation of this approach to validation is that one is unlikely to have a sufficiently extensive set of *in situ* tests to compare models against. Thus, one rarely has a good statistical sample with which to estimate prediction error. Further, we often must extend the use of the models to classes of physical conditions, phenomena or time scales which we have not been able to test in the field. For example, one may wish to use a model which was "validated" for drift inflow to predict large scale regional flow, or we may want to use the flow model as the basis for a transport prediction. Here the prediction error we can calculate does not apply to the problem at hand. Although, there is an expectation that validating a model for a given flow case is a first step in validating a model for other flow conditions or transport, a model validated for inflow is not necessarily a valid model for radically different boundary conditions or different processes.

This brings us to the second component of validation. This approach is commonly called “peer review” or “confidence building.” In this type of validation, the modeling approach is scrutinized. What assumptions were made? Do they make sense? What is the evidence supporting these assumptions? What data were used? What is the sensitivity of the calculation to poor assumptions or inaccuracies in data? Which are the parameters that control the result?

One can prove a model is wrong, but you cannot prove it is right. We can only ask if the model is consistent with observations of all types that were made on the rock mass. Is the model consistent with geologic observations, geophysical measurements, hydraulic measurements, etc.? For example, although the primary prediction is aimed at flux, we can also check to see if the calculated head distribution are consistent with the observed head distribution.

The approach to fracture flow modeling presented here is mindful of both kinds of validation. In Section 5 we explain in detail how prediction error is calculated and we use all the available appropriate data to do this. Below, we present a brief history of our approach to fracture flow modeling which tells how we try to incorporate as much of the physical observations as possible into the model so that we try to insure that the model is consistent with observations a priori. Finally, the physical assumptions made in the numerical code are spelled out in Section 3 and as actual calculations are presented below in Section 7 where we point out what parameters have the most control over the answer we get. Peer review of this process was provided by the Stripa Project Fracture Flows Task Force mentioned in the acknowledgements, whose comments were incorporated into this work.

## 2.0. THE FRACTURE ZONE MODEL

### 2.1. Identification of Fracture Zones with Geophysics

Radar and seismic (acoustic) techniques were used at the SCV site to locate fracture zones. Hydrologic features such as fracture zones represent zones of increased porosity and water content and have different mechanical properties than the matrix rock and these property contrasts should be discernible with geophysics. Geophysical techniques may also find features which are not hydraulically important, so it is important to use the geophysical information in conjunction with a geologic and hydrologic investigation.

Radar signals are sensitive to changes in dielectric constant,  $\epsilon$  and electrical conductivity,  $\sigma$  (Sen, 1981). Radar velocity is a function of  $c/\sqrt{\mu\epsilon}$ , where  $c$  is the electromagnetic wave velocity. The amplitude is a function of  $(\sigma/2)\sqrt{\mu/\epsilon}$ , where  $\mu$  is the electrical permeativity. The values of  $c$  and  $\mu$  are always constant, so a slowness tomogram maps contrasts in  $\sqrt{\epsilon}$  and an attenuation tomogram maps contrasts in  $\sigma/\sqrt{\epsilon}$ .

Both the electrical conductivity,  $\sigma$ , and dielectric constant,  $\epsilon$ , increase with the water content. So, the slowness tomogram should give the most direct correlation with fluid paths in the medium because it depends only on contrasts in  $\epsilon$ . However,  $\sigma$  varies over orders of magnitude whereas  $\epsilon$  only varies by a few percent. As a result, the attenuation tomograms are subject to fewer errors and the image is more clear.

The velocity of acoustic waves in rock are sensitive to changes in rock density, and mechanical properties such as bulk modulus. These are affected by porosity and water content as well as fracture stiffness. The attenuation of acoustic waves is a complex phenomena. Basically, there are two components to attenuation: intrinsic attenuation which is frictional energy loss, and scattering which is due to reflections and refractions at inhomogeneities. Fracture zones increase both types of attenuation of acoustic waves. So, a highly fractured region of the rock is expected to attenuate the seismic waves and also slow them down.

For both radar and seismics, both cross-hole tomographic and reflection techniques were used at Stripa. The tomographic technique uses the direct pulse, or first arrival information. The reflection analysis uses the later arrivals. Thus reflection data provide additional independent information about the fracture zones. Single-hole reflection data are used to locate reflectors in the same manner as common surface reflection surveys. It is possible to determine where the reflectors intersect the borehole and their orientation relative to the boreholes. This information provides a locus of possible reflectors which is a cone passing through this intersection between the boreholes and the reflectors. Use of a directional antenna will make it easier to determine the actual orientation of the feature. A cross-hole reflection technique which is similar to the single-hole method was also used.

An integrated analysis of the radar data was performed by Olsson *et al.* (1988, 1989) which resulted in the identification of major features in the the SCV site. The features were assumed to be planar fracture zones. First, a feature was identified in the tomograms. Then an estimate was made of the location of the intersections between the boreholes and the feature by identifying the feature as a reflector in the single-hole reflection analysis. This gives two possible orientations for the feature, one for each side of the cone intersecting the tomographic plane. If the feature is visible in both the N2-N3-N4 plane and the W1-W2 plane, then there is a further three-dimensional check on the geometry. Then the crosshole reflection data are also checked to see if a unique orientation can be chosen for the zone.

Such analysis was performed separately for both the radar and the seismic data. Then all the data was integrated to obtain four major fracture zones: GA, GB, GH (sub-divided into GHa and GHb) and GI and a minor zone, GC as discussed in Olsson, *et al.* (1989). (n.b.: The "G" in these names stands for geophysics. In the discussion following, we drop the "G" when referring to hydrologic zones. Occasionally, we refer back to zones which were preliminarily identified only with radar data; these are preceded by a "R." The "R" zones are not always located in the same place as the "G" zones of the same name where both radar and seismic data was integrated into the description. However, the hydrologic zones are always in the same location as the "G"

zones.) Table 2.1 summarizes these results with a qualitative classification of the strength of the anomalies associated with the zones made in three classes: S, Strong; M, Medium; and W, Weak. Table 2.1 also includes zone B' which is discussed below.

**Table 2.1. Geophysical Fracture Zones**

Zone	Radar	Seismics	Geoph Logging	Core Logging	Hydrologic
A	S	S	M	M	W
B	S	S	M	M	M
B'	S	S	M	W	S
C	M	S	W	W	W
H	S	S	M	S	S
I	M	W	M	M	S

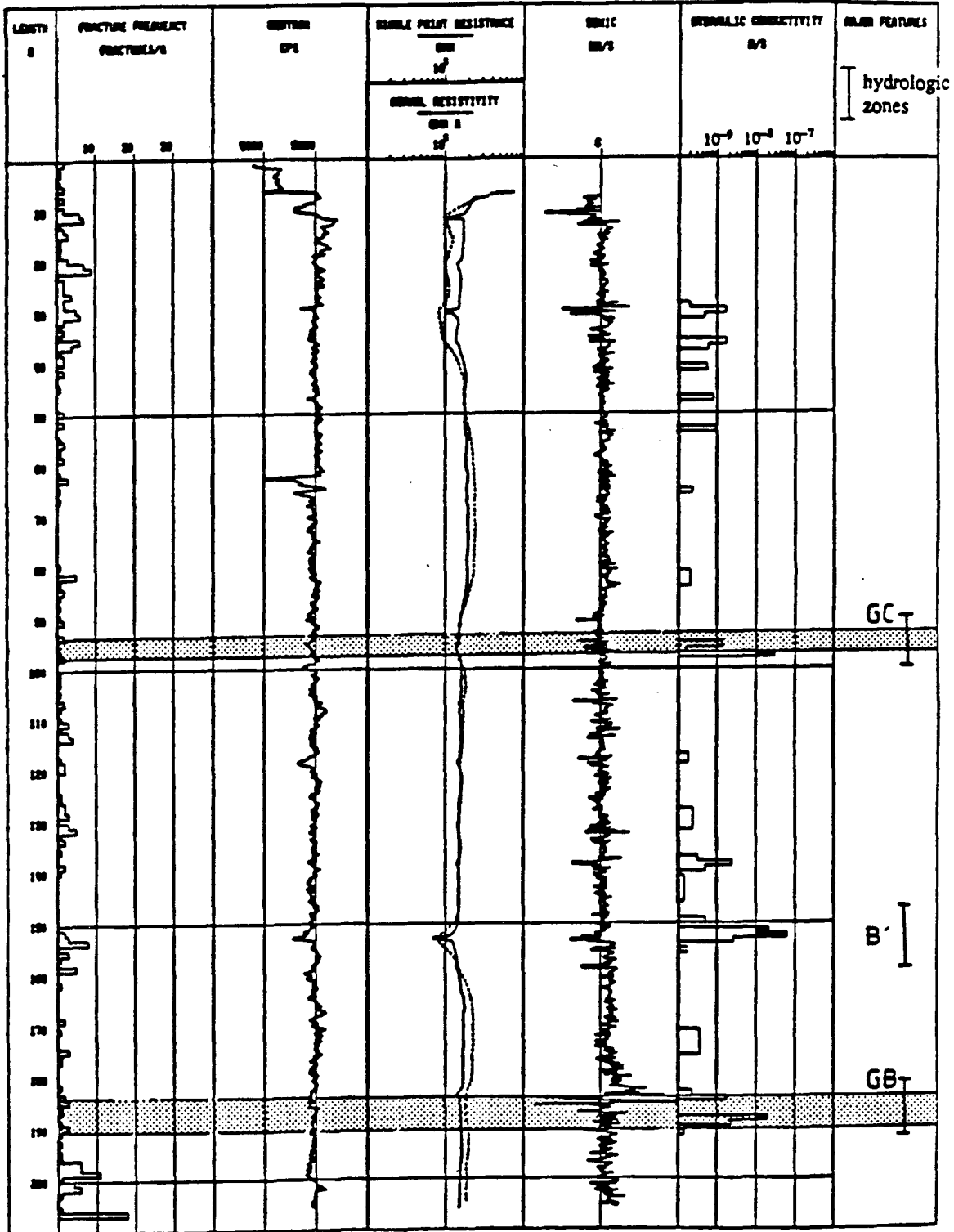
## 2.2. The Hydrologic Conceptual Model

Figures 2.1 (after Olsson *et al.*, 1988) gives an example summary of the borehole data acquired for the N- and W-holes. The geophysical features identified by Olsson *et al.* (1989) are shown on the plots as shaded horizontal bands. Under the hydraulic conductivity column on the right, we have blackened in those conductivities greater than  $10^{-8}$  m/s in order to make the higher conductivity measurements stand out. If the geophysical features are taken as hydrologic zones with a width of about 10 m (denoted by the "I" bars on the right hand sides of Figure 2.1), then they account for about 60% of the measured hydraulic transmissivity measured in the boreholes. Almost all of the remaining 40% of the transmissivity is accounted for in three zones: near 80 m in borehole W2, 152 m in N2 and from 80–90 m in N4. There are strong radar and seismic anomalies at each of these locations as well as anomalies in the geophysical and core logs.

Initially, we tried to account for these hydrologic features by altering the orientations of the five major zones (A, B, C, H and I), an amount consistent with the precision of the geophysical analysis. After determining that no significant improvement was possible, another zone, B', was added to explain previously unaccounted hydrologic anomalies in N4 and N2. Figure 2.2 shows

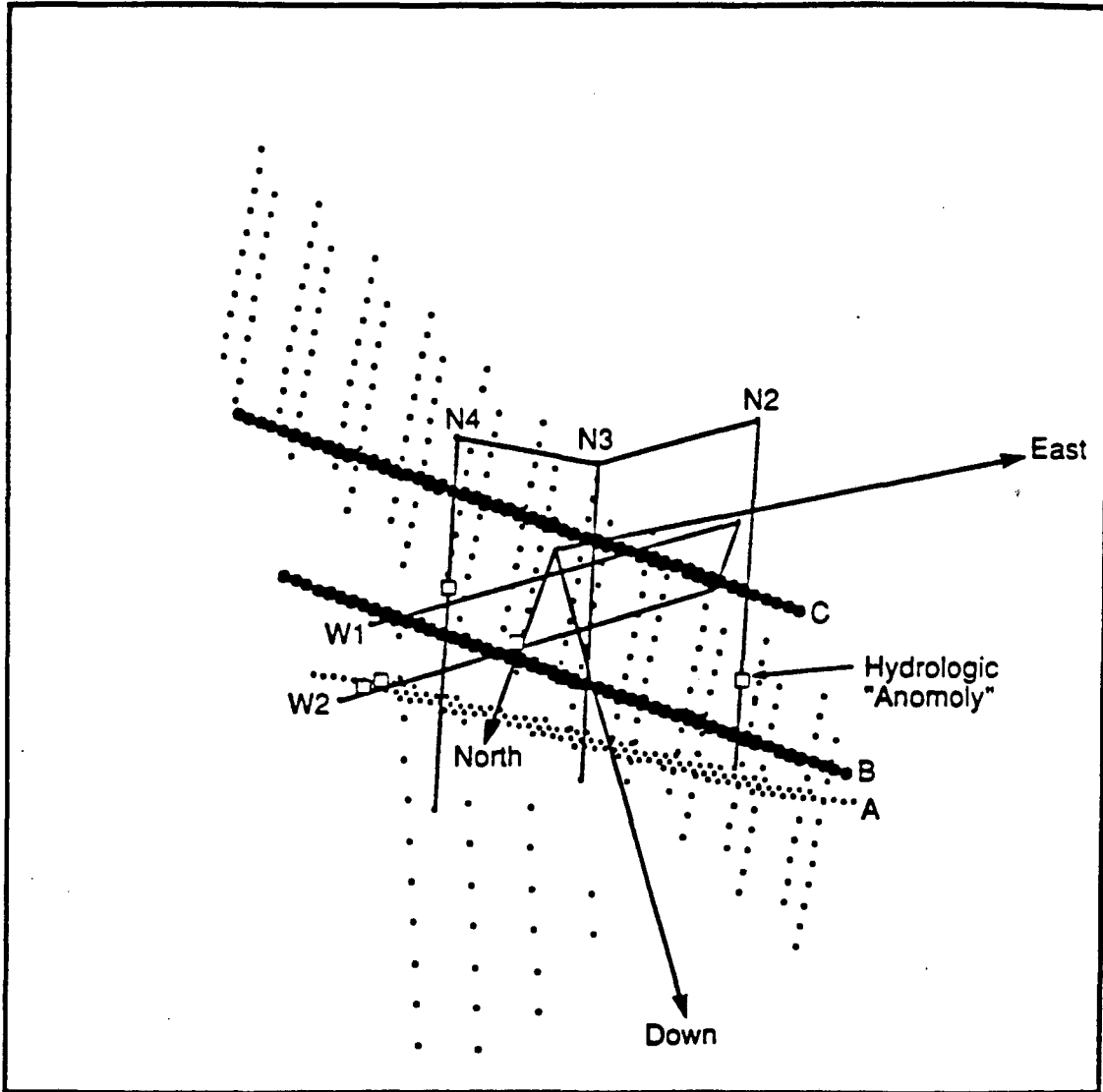


# N2



XBL 903-852

Figure 2.1. Summary data sheet for N2. Hydraulic conductivities greater than 10<sup>-8</sup> m/s have been blackened in. Geophysical features are shown as shaded horizontal bands. Hydrologic zones are marked with "I" bars in the right hand column.



XBL 891-6143

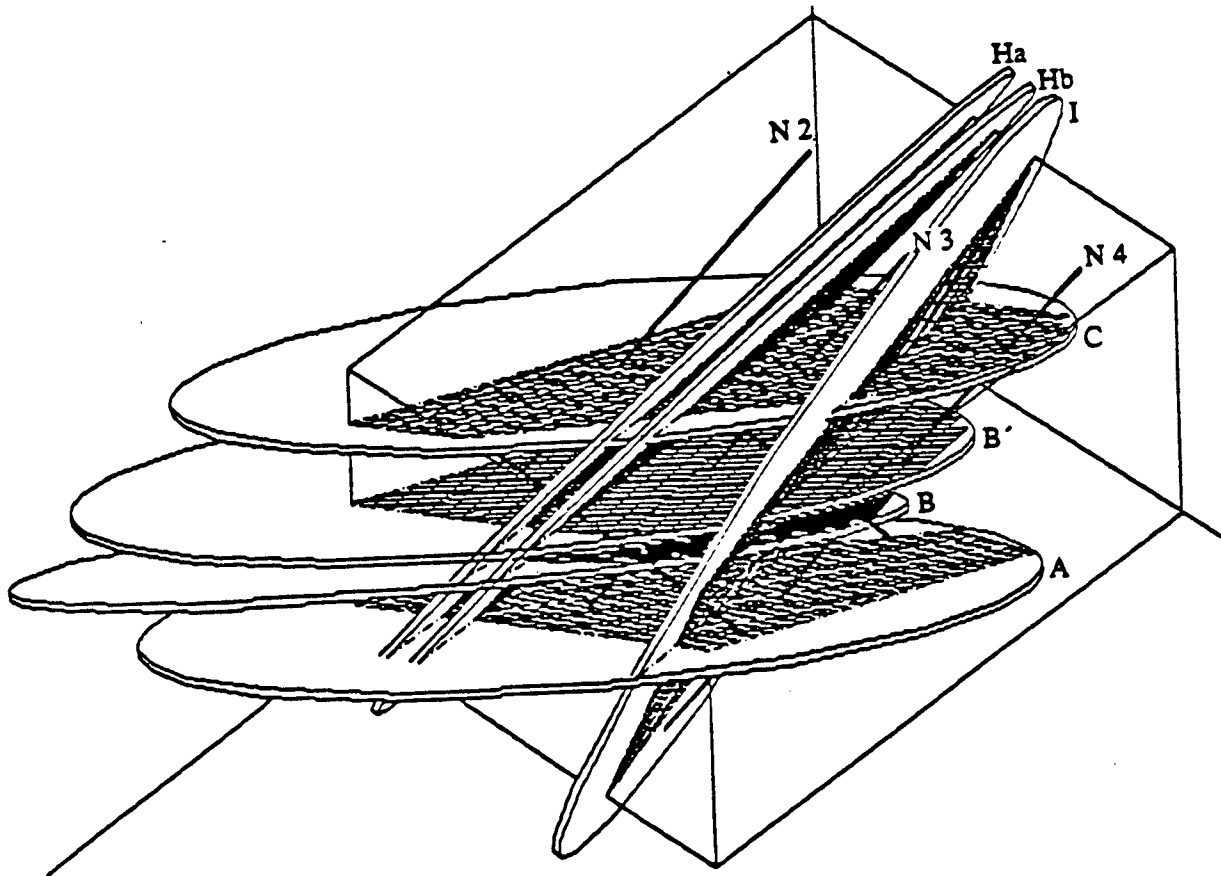
Figure 2.2. A perspective view of the SCV block looking up to the North-East showing zones B and C and two hydrologic anomalies that lie between these zones in a parallel plane.

a perspective plot where zones B and C are represented as dots located on planes. In this figure we are looking along the B and C planes so that the zones appear as dots clustered along a line. In this perspective, one can see that the hydrologic anomalies in N4 and N2 lie on a plane roughly half way between zone B and zone C. For this reason, we chose B' to be a plane between B and C and parallel to the A, B, and C zones. The addition of B' increases the percentage of transmissivity accounted for to about 78%. The resulting hydrologic zone model is shown in Figure 2.3 in a perspective view from the North-West.

The fact that there is no similar hydrologic anomaly where B' would intersect N3 (between N4 and N2) is not a problem for this model. This can easily be accounted for if B' is impermeable in the vicinity of N3.

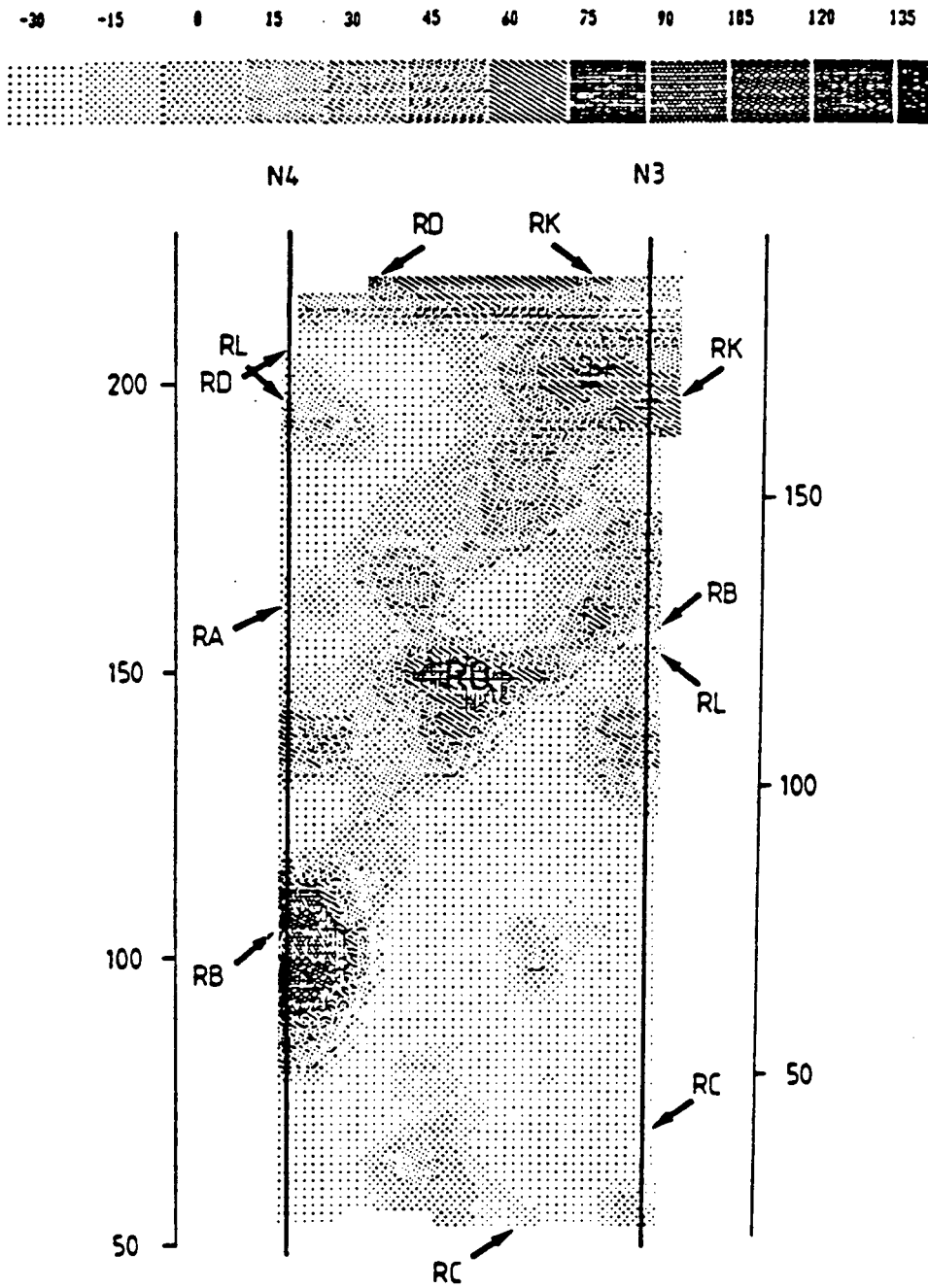
The B' zone also fits in well with some of the geophysical results, but because it did not fit in well with all of the results it was not included in the final geophysical model. The radar results show a very strong, low velocity zone in the section N4-N3 slowness tomograms corresponding to the feature at 90 m down borehole N4 (Figure 2.4, from Olsson *et al.*, 1989) as well as single hole radar reflectors on either side. On Figure 2.4, our B' feature is located at the dark area near N4 labeled "RB" (R for radar). Zone B was originally put in this location when only radar data were considered. Later, the integration of radar and seismic results put zone GB (G for geophysics) up to just below where zone RA is on Figure 2.4. This tomogram shows the B' (or RB) anomaly begins to peter out about 20 m from the borehole. Also, several strong reflectors are observed near the region 152 m down borehole N2. On the other hand, the B' feature is not apparent in any of the attenuation tomograms or the section N2-N4 slowness tomograms, which is why it was not included in the final model. The reason for this absence of B' in the attenuation tomograms is not apparent. However, the N4-N3 tomogram where B' does show up is thought to be more accurate than the N2-N4 tomogram where it does not (O. Olsson, personal communication).

The B' feature may be related to the RQ feature shown on Figure 2.2. It appears that B' may intersect the South edge of the RQ feature. The feature GB then skirts the North edge of RQ



XBL 896-2396

Figure 2.3. The hydrologic zone model shown in perspective from the North-West looking down. Zones A, B, B', C, Ha, Hb and I are shown. Gridding on the planes represents the hydraulic conductors of the template used for annealing.



XBL 896-2397

Figure 2.4. Residual radar slowness tomogram for the borehole section N3-N4 made with a center frequency of 22 MHz.

and produces the largest hydrologic anomaly in N3. It may be that RQ is a step between *en echelon* fracture zones represented by B' and GB. This could also explain the lack of hydrologic activity in N3 at B'. However, it does not explain why B' is again the largest anomaly in N2 unless there is another step in the opposite direction.

One remaining hydrologic anomaly in W2 between zones H and B accounts for 21.7% of the transmissivity. If we allocate this transmissivity partly to H and partly to B, then we can account for 98.7% of the observed transmissivity with a zone model. This makes a certain amount of sense when we consider that the transmissivity measured in the boreholes is not strictly additive. Successive borehole tests actually sample some of the same transmissivity. It is easy to imagine that the high conductivity found between zones H and B in W2 is due to a few conductive features that are related to H and B and possibly related to the intersection of H and B.

The zones in the hydrologic model are represented as disc-shaped planes. As we do not expect the zones to be uniformly permeable, the zones are discretized into flow channels. Any type of discretization could be used. We base the choice of grid with the support of geomechanical investigations of the shear zones explained below. Then, as explained in Chapter 4, we use annealing to find a configuration of active grid elements which can reproduce the observed hydrologic behavior of the system.

### **2.3. Geomechanical Interpretation of the Shear Zones**

Slickenside striations in the two sets of fracture zones indicate that they have undergone shear deformation. Utilizing a database consisting of 3100 logged fractures from borehole core and 900 fractures mapped on the walls of underground drifts, the character of these fracture zones have been analyzed. This data along with numerical fracture mechanics modeling suggest that the higher fluid conductivities in the zones compared with the surrounding rock, may be due to a combination of higher fracture densities, and stresses in the zones that promote dilatancy in fractures with certain orientations. This would also result in anisotropic flow in the zones, with the primary flow direction in the direction of the dilatant fractures.

The H and I fracture zones strike roughly north-south and dip steeply to the east, and the A, B, B', and C zones strike northeast and dip 35 to 40° to the southeast (Olsson *et al.*, 1988). Core logs from these boreholes show that the zones vary in width from 2–15 m, and have a higher fracture density than the surrounding ground (fracture density in the zones varies from 10–30 fractures/m, compared with 1–20 fractures/m in the ground surrounding the zones). The borehole data base contains information on 3100 fractures of known orientation. The five boreholes (N2, N3, N4, W1, W2) intersect the fracture zones in over 20 locations. Scanline surveys were taken in the drifts surrounding the SCV site, which contain photographs, orientations, and associated information for 900 mapped fractures. In addition, a few detailed wall maps were made along some of the underground drifts. This information is used in the context of the overall geological framework of the area surrounding the Stripa mine, and stress measurements indicating the present stress state (Chan *et al.*, 1981; Carlsten, 1985; Olsson *et al.*, 1988).

Based on an analysis of this data, there is evidence that the fracture zones in the SCV block have undergone extensive shear deformation. Slickensided fractures in the zones indicate that these zones contain faults. Also, where these zones intersect underground drifts, cracks that splay from slickensided fractures can be observed. We infer that the zones identified by radar and seismic tomography are mostly fault zones. Certain patterns of fracturing can be expected in fault zones in brittle rock (see Section 2.3.2 below), and some of these patterns have been observed in the SCV block. This data, along with numerical modeling suggest that the higher fluid conductivities in the zones compared with the surrounding rock may be due to a combination of higher fracture densities, and stresses in the zones that promote dilatancy in fractures with certain orientations.

Laboratory tests of intact rock under direct shear by Tchalenko (1970) and others indicate that initially a set of *en echelon* fractures form, finally linking up into a throughgoing fault. Shear fractures formed in this way can show a remarkable resemblance to faults mapped in the field (Tchalenko, 1970; Sibson, 1986). On the other hand, the field studies of Segall and Pollard (1983), Martel *et al.* (1988), and others indicate that faults can form from slip along pre-existing

joints. As slip increases along these pre-existing fractures, secondary fractures form and link the slipping fractures to form throughgoing faults.

In both of these scenarios, a high density of fracturing develops adjacent to the throughgoing slip surfaces. These associated fractures can be subjected to stress states much different than fractures in the surrounding rock. For instance, *in situ* stresses in the ground are normally compressive in all three principal directions. However, even under compressive *in situ* stresses, slip along fractures can locally induce tensional stresses (Rodgers, 1980; Segall and Pollard, 1980; Sibson, 1986). Fractures that open in these extensional regions can have hydraulic conductivities much greater than fractures in surrounding areas which are under compressive stresses in all directions. Mineralized areas commonly occur in regions such as these (Sibson, 1981). Also, extensional fractures in shear zones and faults will have a preferred orientation, and this can result in an anisotropic flow regime in the shear zone. Thus the orientation and density of fractures in a rock mass is not sufficient to determine flow properties. It is also necessary to consider the presence of shear zones or faults, and the overall *in situ* stress state.

### 2.3.1. Fractures in the SCV Block

Before looking at the fracture zones themselves, we give a general overview of the pattern of fracturing in the SCV block. The granite hosting the fracture zones at the SCV site is fractured itself, albeit not as extensively as the fracture zones. Over 3000 fractures are intersected by the five boreholes in the SCV block. A statistical analysis of the orientations and locations of these fractures is given in Olsson *et al.* (1988). When fracture data from all five boreholes are combined, three main fracture orientation clusters are observed. Two of these roughly coincide with the orientations for the fracture zones shown in Figure 2.3, and the third cluster group is sub-horizontal. The sub-horizontal fractures represent a small percentage of the total number of logged fractures, but this is partly due to the low plunge of the boreholes drilled through the SCV block. Next, we focus on the fracture zones.

The NE-striking, low dipping fracture zones includes A, B, B', and C. We focus on zone B', since it represents a highly conductive feature in the SCV block. Poles to the fractures in zone B'



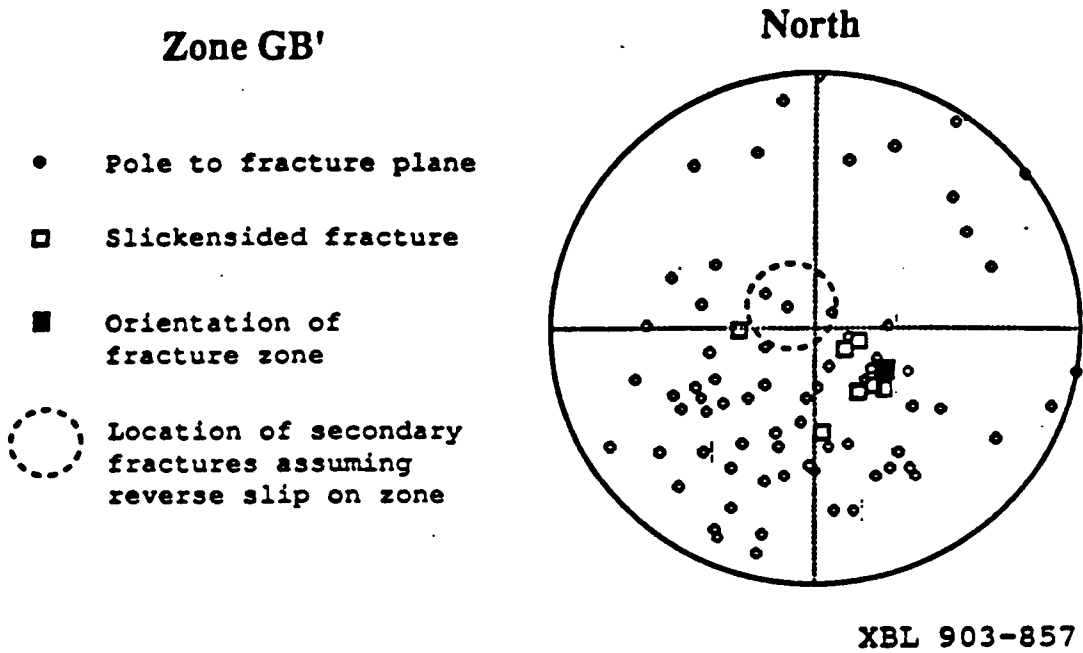


Figure 2.5. Poles to the fractures in zone GB' (upper hemisphere) where it intersects borehole N4 (depths 80-95 m).

where it intersects borehole N4 (depths 80–95 m) are shown in Figure 2.5. The dark square in Figure 2.5 represents the best estimate for the orientation of this zone, based on where it intersects boreholes and as described above. Some of the fractures in this zone are slickensided, and these fractures are marked with hollow squares in Figure 2.5. Figure 2.5 shows that most of the slickensided fractures and many of the non-slickensided fractures are sub-parallel to the orientation of the zone. We infer that the slickensided fractures represent major faults that parallel the zone as a whole. Steeply dipping striae on slickensided fractures that are sub-parallel to the zone indicate a dip-slip type of motion for the northeast striking, low dipping zones. Sub-horizontal cracks that splay from some of these slickensided fractures can be seen on the drift walls and indicate a reverse dip-slip motion (Petit, 1987). This sense of motion is consistent with stress measurements in the SCV site, indicating a present maximum horizontal compression oriented northwest (Chan *et al.*, 1981; Olsson *et al.*, 1988).

Many, if not most of the fractures logged where zone B' intersects borehole N4 may not be associated with the shear zone, but may instead represent "background" fractures in the host rock (as described in the previous section). Certainly the fractures that are sub-parallel to the zone are more likely to be related to the zone. Also, secondary extensile fractures may be associated with slip across the zone. Secondary extensile fractures are apparent in the driftwalls, but are difficult to identify from borehole logs. These dilatant fractures should have a narrow range of orientations, related to the orientation of the zone, the direction of slip in the zone, and the orientation of the far field stresses (e.g., Petit, 1987). For the northeast-striking, low dipping zones under reverse dip-slip motion, the fractures should plot within the circle shown in Figure 2.5. Thus fractures within this circle may be extensile fractures that are associated with reverse-slip on this zone. Also, fractures with this orientation may be contributing more to the overall conductivity of the zone than the fractures that are sub-parallel to the zone. For instance, Teufel (1987) shows that due to the accumulation of gouge, the permeability of fractures under shear in the laboratory decreases with increasing shear displacement. Thus the slickenside fractures may not be contributing substantially to the conductivity of the zone. Teufel (1987) notes, however,

that the rock immediately adjacent to the sheared fracture may be highly conductive due to extensile microfracturing formed from the shearing.

Fractures within the circle in Figure 2.5 are sub-horizontal. Based on this, we infer that sub-horizontal fractures would be preferred pathways for flow within the NE-striking zones in the SCV block. However, in Figure 2.5 only a few fractures occur with this orientation. This may be partly due to the bias against sub-horizontal fractures appearing in a borehole that plunges only 18.6°. Sub-horizontal fractures have been mapped in the driftwalls surrounding the SCV block, and some of these fractures have been identified as secondary fractures due to slip in the NE-striking zones.

The N-S striking, steeply dipping zones includes H (Ha plus Hb) and I. We focus on the H zone, since it is the most conductive feature in the SCV block. Poles to the fractures in zone H where it intersects borehole W1 (depths 48–63 m) are shown in Figure 2.6. Steeply dipping striae on slickensided fractures, along with shallowly-dipping splays suggest reverse dip-slip movement along zone H. Secondary fractures due to reverse slip on this zone should strike N-S and dip 10–40° to the east, as shown in the circle in Figure 2.6. Because boreholes W1 and W2 dip only 5°, there will be a bias in the boreholes against fractures with this orientation. However, shallowly dipping fractures associated with zone H are seen in the driftwalls. We infer that shallowly dipping (to the east), N-S striking fractures would be preferred pathways for flow in these zones.

### 2.3.2. Geomechanical Modeling of the Shear Zones

The possible high conductivity of secondary fractures in the zones, compared with slickensided fractures in the zones and fractures outside the zones is supported by the results of the numerical example shown in Figure 2.7. We have considered a simplified two dimensional elastic model under far field compressive horizontal and vertical stresses  $\sigma_1$  and  $\sigma_2$ . According to Chan *et al.* (1981) and others, the maximum principal *in situ* stress is horizontal with a magnitude of approximately 24 MPa on the 360 m level. This stress is oriented northwest and therefore perpendicular to the NE-striking zones. The minimum principal stress is vertical with a magnitude of

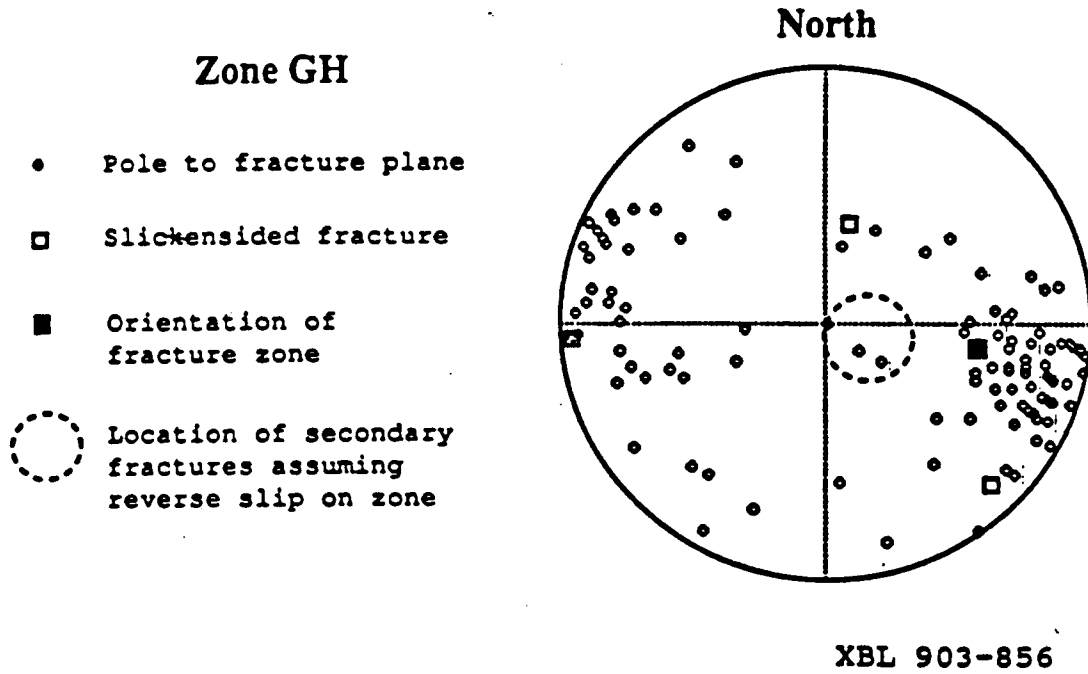
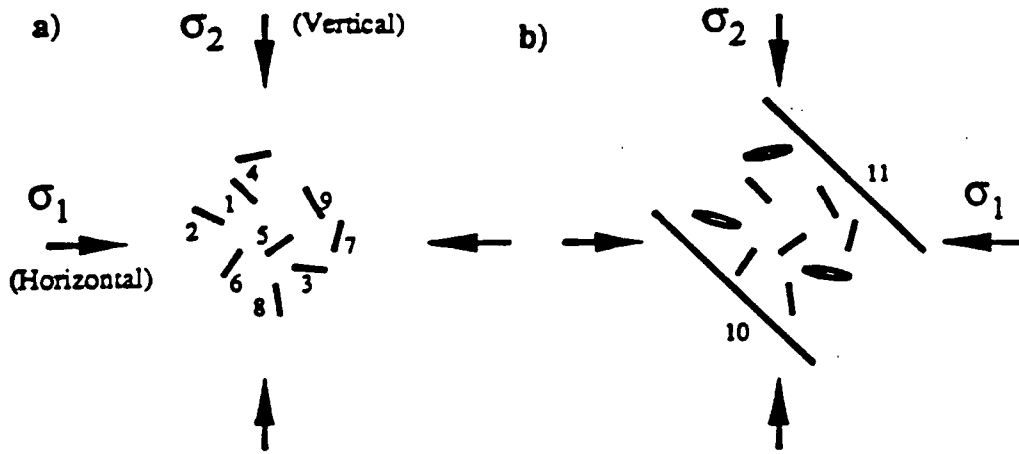


Figure 2.6. Poles to the fractures in zone GH (upper hemisphere where it intersects borehole W1 (depths 48-63 m)).



a)				b)			
#	$K_I$	( $\text{MPa}\sqrt{\text{m}}$ )	$K_{II}$	#	$K_I$	( $\text{MPa}\sqrt{\text{m}}$ )	$K_{II}$
1	0		-7.8	1	0		-5.3
2	0		-4.4	2	0.92		-4.5
3	0		3.7	3	0.55		0.4
4	0		5.4	4	0.05		7.8
5	0		8.1	5	0		12.9
6	0		5.9	6	0		15.5
7	0		2.3	7	0		12.4
8	0		-4.4	8	0		5.3
9	0		-7.5	9	0		-0.6
10	0		-	10	0		-38.6
11	0		-	11	0		-38.6

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Figure 2.7. (a) Nine randomly distributed fractures subjected to the maximum horizontal and vertical stresses on the 360 m level; (b) Two long fractures representing the slickensided fractures in the NE-striking zones are added. The table on the figure lists average Modes I and II stress intensity factors for each crack for the two cases.

of approximately 9 MPa on the 360 m level. In Figure 2.7a, nine randomly distributed fractures are subjected to these stresses. For simplicity it is assumed that the fractures do not cross each other, and each of these fractures have been given a length of one meter. In Figure 2.7b we look at the same system of fractures, except now two long fractures representing the throughgoing slickensided fractures that dip  $40^\circ$  are added. The long fractures each have a length of 16 m and are separated by 7 m. Thus in Figure 2.7b the nine fractures are subjected to the stress state within one of the NE-striking, low dipping zones i.e., the A, B, B' and C zones.

Due to the application of the *in situ* stresses, the modes I and II crack-tip stress intensity factors,  $K_I$  and  $K_{II}$ , respectively, have been calculated at each of the crack tips utilizing a numerical algorithm based on the work of Kachanov (1987). This algorithm has been modified to account for cracks that are closed by the compressive stresses but can still shear (frictionless surfaces are assumed for closed cracks). The  $K_I$  and  $K_{II}$  for each crack is presented in the table below Figure 2.7. Each crack has two crack tips, and in Figure 2.7 we only show the average of the two crack tips.  $K_I$  is an indication of the opening of the crack, and  $K_{II}$  is an indication of the shearing of the crack (Lawn and Wilshaw, 1975).  $K_{II}$  can be both negative and positive indicating shear in one direction or the other, while  $K_I$  can be either positive or zero.  $K_I = 0$  indicates that the crack is closed.

Figures 2.7a and b demonstrate the importance of shear zones in localizing flow in the rock mass. In Figure 2.7a, each of the fractures are under different amounts of shear but they are all closed ( $K_I = 0$ ). In Figure 2.7b, however, due to the localization of shear from the long slickensided fractures, fractures with certain orientations have a positive  $K_I$ , and are therefore open. These fractures are sub-horizontal, which agrees with the expected orientations as shown in Figures 2.5 and 2.5.

Figure 2.7 also gives the stress intensity factors for the long fractures, and they remain closed. The sign of  $K_{II}$  for the long fractures indicates a reverse dip-slip motion, as the field data suggests for the NE-striking zones. Thus this numerical example supports the idea that the conductivity is greater in dilatant fractures in the zones rather than the main throughgoing fractures

that are sub-parallel to the zone.

Figure 2.7 also points out a technical difference between a fault zone and a fracture zone. A fracture zone may contain a high density of fractures, but under compressive far field stresses, the fractures will be closed, as in Figure 2.7a. A fault zone, on the other hand, by containing dilatant fractures, can be more conductive, even if the density of fractures is lower.

## 2.4. Conclusions

Seismic and radar tomography of the SCV underground test area in Stripa, Sweden have detected two major sets of fracture zones. We have evidence that they are fault zones under reverse dip-slip motion. Associated with slip in the zones, secondary fracturing has been observed. For the NE-striking, low dipping zones, the secondary fractures are sub-horizontal, and for the N-S striking, steeply dipping zones, the secondary fractures strike N-S and dip 10–40° to the east. Numerical modeling indicates that under the present stress state in the SCV block, the sub-horizontal secondary fractures could be open and have a much higher conductivity than other fractures in the SCV block. This, along with the higher fracture densities in the zones, may explain why the conductivity in the zones is greater than the surrounding ground. Also, this will cause anisotropy in flow in the zones with preferred pathways in the direction of the secondary fractures. Though not discussed in this paper, there is evidence that zone intersections may be important in controlling flow through the SCV block. Future work looking at the properties of zone intersections is recommended.

Based on this work, we chose a zone model where the zones are modeled as planes. The planes are discretized using a square grid of conductors to form the possible paths for fluid flow. The grid elements are constructed along strike and dip lines. In this way we allow for the horizontal conductors indicated by the geomechanical observations. As a first try, each grid element is assumed to have the same conductance, so if the geomechanical observations are correct, we expect that fewer of the dip direction elements will be active, i.e. permeable than those in the strike direction.

In order to determine which of the grid elements are active and which are inactive, we can employ a technique called "Simulated Annealing." Simulated annealing is an inverse technique which is used to find the pattern of conductances which cause the model to behave in the way the the insitu tests behaved. First, well tests that were performed insitu are simulated in the model using the network generator, CHANGE, plus the fracture flow code, TRINET. These numerical models are described in Section 3. Then the annealing process (Section 4) is used to eliminate or restore a conducting grid element and the response to the well test is recalculated with TRINET. We compare the behavior of the new model to the old and decide whether or not to keep the change. This process is repeated many times until the model matches the observed behavior. The algorithm is designed to find an arrangement of conductors that responds to the well test in nearly the same way that the real well test was observed to respond. This final configuration of the grid can be considered as an equivalent "rastorized" version of a possible conductance geometry. Finally, the boundary conditions imposed on the model can be changed in order to make predictions of flow that have not been measured.



### 3.0. THE NUMERICAL MODEL

#### 3.1. The CHANGE Model

The channel network generator, CHANGE (Billaux, *et al.*, 1988), is used to construct a finite element mesh for the hydrologic conceptual model. CHANGE can generate a stochastic network of conductors or any regular grid of conductors. In the finite element analysis, each element is assumed to represent a one-dimensional conductor, i.e. a pipe. For the purpose of calculating flow, we do not need to describe the shape of the conductor. We only need to know the location of the endpoints and the conductance of the element which has the units of permeability times cross-sectional area:  $m^3/s$ . An element is not necessarily meant to precisely represent a specific channel. It may represent the equivalent conductance of several channels combined.

#### 3.2. TRINET

TRINET (Karasaki, 1988) is used to calculate the hydrologic response of the zone model to the simulated well test. TRINET is a finite element code for simulating fluid flow and advective and dispersive solute transport in 3-dimensional networks of one-dimensional conductors. The code incorporates a mixed Lagrangian and Eulerian scheme with adaptive gridding for transport calculations. The model avoids numerical dispersion by creating new Eulerian grid points instead of interpolating the advected profile back to the fixed Eulerian grid.

The code first solves the flow field using a simple Galerkin finite element method. The flow can be either steady-state or transient, where the time derivative is treated in a usual finite difference manner. From the pressure distribution at a given time the velocity distribution in the fracture network is calculated. Since linear shape functions are used to solve the flow field, the velocity is uniform within a given element. Elements are sub-divided as necessary for transient calculations. The advection-dispersion equation for mass concentration is then solved by decoupling the equation into two stages. In this way it is possible to solve the advection term indepen-

dently of the diffusion term and minimize the numerical dispersion.

First, the advection equation is solved by using the method of characteristics. The concentration profile at the end of the calculation in the previous time step is, in effect, the initial-value distribution for the new advection problem. This profile is advected explicitly in the Lagrangian manner according to the velocity in each element. The advected front is placed on the Eulerian grid and new nodal points are generated to preserve the exact shape of the front. At every time step the element catalog is revised and the nodal points are renumbered to keep the band-width minimized. Because it is necessary to preserve the geometry of the fracture network itself, the original nodal points must be kept intact. The method of single-step backward particle tracking or reverse streaklines is used to obtain the concentration values for these fixed nodal points.

The new concentration profile at the end of the advection stage is now the initial-value for the dispersion calculation in the second stage. The dispersion is treated in the usual finite element manner.

In the present study, only the flow is simulated using TRINET. In this case, we need to know more about the structure of the conductance elements. For the purpose of flow, it does not matter whether we have a high conductivity channel with a small cross-sectional area or vice versa. For transport, the fluid velocity through the high conductivity channel with a small cross-sectional area will be larger than for the opposite case. This means, not surprisingly, that if we construct a model that predicts flow, we do not have enough information to predict transport. Independent information would have to be used to relate flux to velocity in each element.

### 3.3. Scale of Analysis (“Resolution”)

In modeling the hydrology at the SCV site, we opted to use a channel network model as was described earlier. Thus, the system will be represented by a finite number of elements. It is now necessary to determine the scale of analysis, i.e., how finely we will discretize each zone. The scale of analysis is constrained by the size of the physical memory of the computer. This effectively sets the maximum number of conductors that can be modeled which in turn deter-

mines the maximum resolution. With the current configuration of the computer at LBL, the maximum number of nodes that is feasible for the annealing process is about 10,000. This is roughly equivalent to having a 10 m grid spacing on each fracture zone.

Another factor that affects the scale of analysis is the density of the monitoring points. The minimum spacing of the conductors should be less than that of the monitoring points. One would like to have more than a single conductor between two monitoring points to reconstruct the geometry of the fracture zone. This effectively sets the minimum size of the mesh. It is in general desirable to have as many monitoring points as possible to accurately characterize a site. In the case of SCV site, the average spacing of monitoring points is on the order of a few to several tens of meters. The grid spacing needs to be smaller than the distance between monitoring points, which again gives approximately 10 m as the appropriate grid spacing.

Finally, the purpose of our simulation is to resolve an equivalent hydrologic geometry of the fracture zones and not individual fractures. It is known that the drawdown effect due to open W2 is felt all across the site to R-holes at the ventilation drift. The scale of the fracture zones, therefore, is estimated to be on the order of several tens of meters to a few hundred meters in extent. From the above point of view, a grid spacing of about 10 m is again reasonable. In order to divide the plane by an even number, the spacing that was actually used placed a grid of 8.8 m squares on each plane.

### **3.4. Treatment of the D-Holes as One Big Hole**

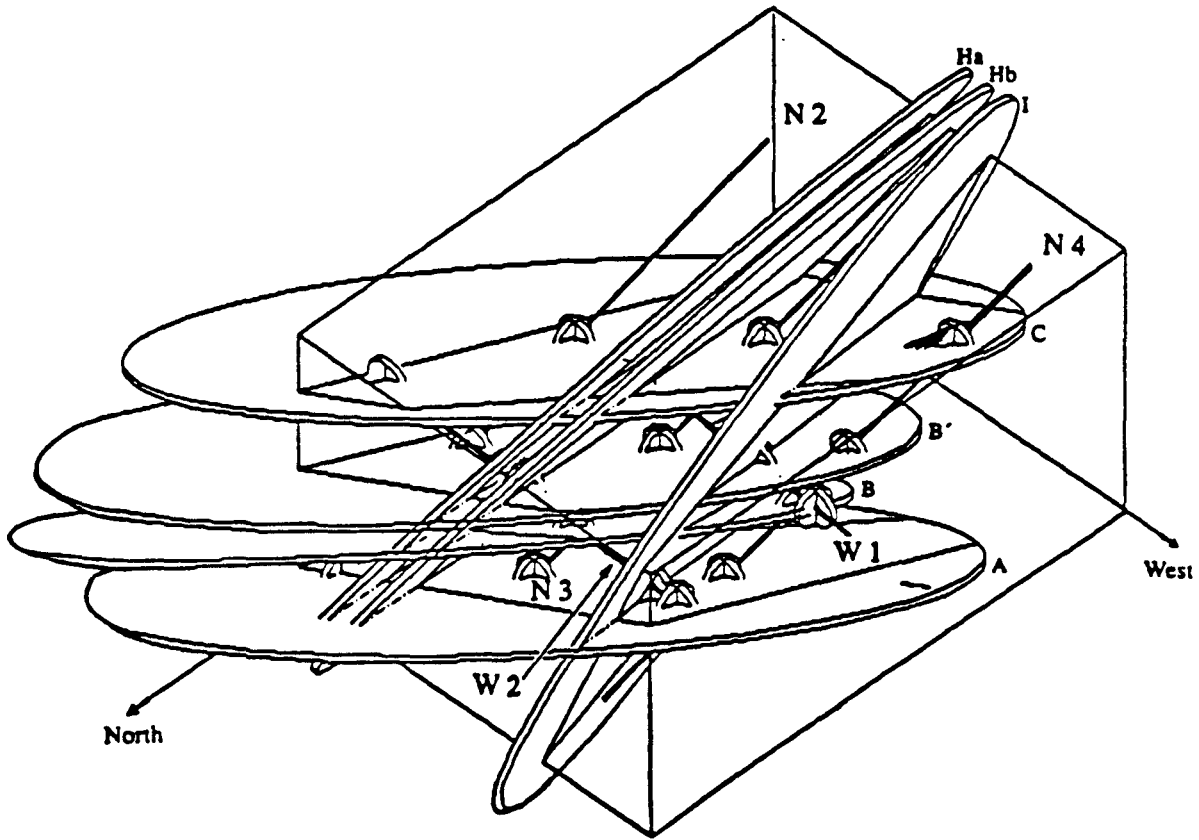
The current resolution of the analysis is on the order of 10 m, but the five D-holes are only about 0.1 m in diameter and 1.2 m apart. In predicting the inflow into the D-holes, therefore, they have to be treated as one big hole.

Because the background permeability is neglected, the predicted inflow into the drift will be zero except where the zones intersect the drift. This may be a good assumption based on the fact that the background permeability is a very small percentage of the zone permeability. One can find analogous cases in any one of the N and W holes, where nearly all of the transmissivity

is localized in a few zones.

### **3.5. Connecting the Zones to the Boreholes**

As the boreholes in general are about 0.1 m in diameter, it is extremely unlikely that any of the boreholes will intersect a channel in the zones. To insure a hydraulic connection between the zones and the wells, small "fins" were added to the boreholes at the points where they intersect the zones. These fins are made of two orthogonal discs 6 m in diameter. The construction of the fins allowed the intersections between the fin discs to be channels which in turn intersect zone channels becoming connections to the borehole. The fins and zones are shown in Figure 3.1.



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Figure 3.1. The hydrologic zone model with the fins that connect the zones to the boreholes.

#### 4.0. SIMULATED ANNEALING

LBL has been developing an inversion technique called "Simulated Annealing" which can be used to construct a system which is functionally equivalent to the observed system: i.e., a model which simulates the same behavior as the observations we have. We describe here how to use annealing to find an equivalent fracture network model. The fracture network model is "annealed" by continually modifying the base model, or "template" such that the modified systems behave more and more like the observed system.

Hydrologic inversion models developed in the past, such as the conjugate gradient method, or maximum likelihood method (Carrera and Neuman, 1986), were focused on determining the conductivity values when the pattern of conductors is known or when everything is well interconnected as in the porous medium case. Annealing could theoretically be used to do this type of inversion, but might be relatively inefficient in this role. On the other hand, these porous medium techniques work poorly when one needs to completely turn off the conductivity of a portion of the region. Thus they are not the technique of choice for fracture systems when we wish to determine how the conductive features are connected.

Annealing is specifically designed to determine an appropriate pattern of conductors among a set of possible configurations. In fracture hydrology, we think that the pattern of conductors is responsible for the first order behavior. In other words, it is most important to know how the system is connected. Annealing is designed to find connections equivalent to those in the field. This is achieved by methodically searching patterns to see which ones behave like the ones observed in the field. Simulated annealing is actually the statistical technique which controls the acceptance or rejection of trial modifications. The job of simulated annealing is to find near optimal solutions of complicated function with many possible solutions and local minima.

The set of possible configurations is based on a template or base model which specifies all of the possible connectors. We are pursuing two different approaches to defining the template.

The first approach has been described above. We make the assumption that the behavior is controlled by fracture zones, so we only allow connectors to exist within the zones. This approach has the advantage that it is efficient because it uses the information gained from geophysics and geology *a priori*. The second approach is to use a three-dimensional regular grid of conductors as the initial template. This approach is described in Appendix A and has the advantage that the hydrologic responses drive the result more directly, but the allocation of conductors may be inefficient.

For either approach, the resolved pattern is not guaranteed to be the same as the one in field, and for this reason we call it an equivalent system. The equivalent systems are non-unique. This means we can find a range of systems which behave the same way the real system behaves. The range of systems can then be used to make a range of predictions.

Once an equivalent pattern has been determined, we can compare the behavior of the model to that of the observations. In some cases, it may be that pattern alone does not allow one to create a sufficiently accurate model. One could refine the template in the vicinity of the poor match in order to provide for greater contrasts in conductance. However, for very large contrasts one might wish to use annealing to discern the variability of conductance among elements. This has been observed in one case (Davey *et al.*, 1990).

The process of annealing is summarized as follows. *In situ* well tests are simulated on the model using TRINET. We then define the "energy,"  $Q_C$  of our current model configuration, (referred to as  $C$ ), as the squared sum of the differences between the measured and simulated heads, taken at a set of locations at discrete time intervals. Next we change the model by switching a randomly chosen channel "on" (i.e. conducting) if it is "off" (i.e. non conducting) or vice versa and then repeat the well test simulation. If the changed model gives simulated well test data closer to the real measured values, i.e.  $Q_C$  is decreased, then the changed model is kept. However, if  $Q_C$  is increased by the change, then the change will be kept with a certain probability,  $P$ , which is a function of a weighting factor,  $T$  (called the "temperature"), and  $Q_C$ . The temperature is decreased as the number of iterations increases to make it more and more unlikely that

an unfavorable change will be accepted. At first, a high value of  $T$  allows the algorithm to jump up out of local minima and continue searching for a better region of the function. Later, lowering the temperature tends to confine the search for a minima, so the algorithm can converge.

#### 4.1. Annealing Theory

This optimization technique employs an analogy between optimizing a function with many local minima and the process of annealing a metal to reach its lowest energy state. Thus, simulated annealing can be viewed as a process of minimizing an energy function over a set of possible configurations of elements.

Simulated annealing has been defined in analogy with the thermodynamic process of cooling or annealing of a metal (Kilpatrick, *et al.*, 1983; Tarantola, 1987). A system composed of atoms of a metal in thermal equilibrium at a temperature  $K$  has a range of possible energies depending on the configuration. Thermodynamically, low energy states are more likely, but at any temperature there is still some chance of being in a high energy state. Metropolis *et al.* (1953) used this conception of energy states to simulate changes in a thermodynamic system.

To use simulated annealing on a general problem, one needs a set of possible configurations, a way of randomly changing the configurations, a function one would like to minimize, and an annealing schedule of temperature changes (Press, *et al.*, 1986).

We can define the set of all possible configurations using our base model:  $M$ , i.e. the template which is the set of all possible pipes or channels. The channels have two possible states: they are either off or on. The set of all possible configurations is the set of all combinations of off and on pipes. Let  $C = (C_m, m = 1 \dots M)$  denote a configuration of on and off pipes, where  $C_m$  is a binary random variable associated with each pipe.

We now change the system. First, we use some probability function to randomly select a pipe. If the pipe is on, we turn it off, and if the pipe is off, we turn it on. In this way we find a new configuration which is in the neighborhood of  $C$  where the neighborhood of  $C$  is all configurations one step away from  $C$ . Let  $C$  be the configuration at iteration  $n$  and  $G_C$  be its



neighborhood. When we anneal the system, we randomly select a configuration  $C'$  from  $G_C$  at each iteration  $n$ , and compare the two energy functions.

The energy functions we use are a measure of the difference between the observed and the simulated system response. We consider energy functions of the form:

$$Q = \sum_j |f(o_j) - f(s_j)|^m \quad (4.1)$$

where

$o_j$  = a vector of observed responses

$s_j$  = a vector of simulated responses

$m$  = a constant, and

$f(\cdot)$  is a real monotone function.

The observed measurements could be hydrologic, geological, or geophysical.

The energy function, scaled by the temperature, is used to decide whether the system should make a transition to a new configuration. The temperature is lowered as the algorithm progresses, to make it increasingly unlikely that a transition to a higher energy state will occur.

We assume a probability distribution on the configurations can be expressed as a Gibb's Distribution:

$$P(C) = \frac{1}{k} e^{-\left[\frac{Q(C)}{T}\right]} \quad (4.2)$$

Where  $k$  is a normalizing constant which assures that the sum of the probabilities of all possible configurations is unity. We assume that this constant exists, but it is very difficult to evaluate because we must know the energy for every possible configuration to compute  $k$ . Therefore, we can not compute the absolute probability of any given configuration because we do not know  $k$ . However, we can compute the relative probability of any given configurations. For example, we could say that a configuration would be twice as probable as another.

Further, we know that if the probability function is a Gibb's distribution, then this is equivalent to modeling  $C$ , the current configuration as a Markov Random Field. A Markov Random Field exists on a neighborhood system,  $G$ , if the probability defined meets two conditions. The first condition is that the probability of selecting any configuration in the system is greater than zero. The second is that the probability of making a transition from  $C$  to  $C'$  depends on  $C$  and  $C'$ , and whether  $C'$  is in the neighborhood of  $C$ . Past history, such as the configuration we selected before  $C$ , does not tell us anything about the probability of moving from  $C$  to  $C'$ . Therefore, the probability of moving from one configuration to another can change with the iteration, but it does not depend on which configurations have been examined in the past. This means we can examine a series of configurations without remembering how we moved from one to the next and we can still compute the relative probability of each configuration.

At each iteration  $k$ , given  $C$ ,  $G_C$ , the neighborhood, and  $T$ , the temperature, we can find a matrix of transition probabilities. The probability we will move from configuration  $C$  to  $C'$ , given our current configuration  $C$ , is equal to the probability that we select  $C'$  to compare with  $C$  multiplied by the probability that the system would make the transition to a given  $C'$ . That is:

$$P\{C \rightarrow C' | C\} = \begin{cases} 0 & \text{if } C' \notin G_C \\ P(C' | C) \cdot 1 & \text{if } C' \in G_C, C' \neq C \\ & Q(C') - Q(C) \leq 0 \\ P(C' | C) \cdot e^{-\left[\frac{Q(C') - Q(C)}{T}\right]} & \text{if } C' \in G_C, C' \neq C \\ & Q(C') - Q(C) > 0 \end{cases} \quad (5)$$

and the probability of not accepting the change to  $C'$  is:

$$P(C \rightarrow C | C) = P(C \rightarrow C) = 1 - \sum_{\{C': Q(C') \leq Q(C)\}} P(C' | C) - \sum_{\{C': Q(C') > Q(C)\}} P(C' | C) \cdot e^{-\left[\frac{Q(C') - Q(C)}{T}\right]} \quad (4.3)$$

It remains to discuss the temperature schedule which is to lower the temperature as annealing progresses. Physically, this means that as annealing progresses we are less and less likely to keep changes which increase the energy of the system. At this time, there is a theory which relates the temperature schedule to the convergence properties of annealing. This theory (Hajek, 1988) shows that a temperature schedule which is inversely proportional to the log of the iteration number will converge in probability to a set of minimum energy states.

However, a temperature schedule of this type is prohibitively slow. Furthermore, we do not necessarily want to find the minimum energy configuration. We are only interested in finding several good solutions, not in certain convergence in probability to a set of minimum energy states. Therefore, use of Hajek's temperature schedule is over constraining for our purpose.

The temperature schedule we use here is only justified heuristically: it works. We have followed the suggestion of Press, *et al.* (1986) and decreased the temperature after a number of iterations which is sufficient to result in accepting 50 changes. Each interval of the schedule with constant temperature is called a step. At the end of each iteration,  $k$ , the temperature,  $T_k$ , is decreased using a geometric series,

$$T_{k+1} = T_k t^k \quad (4.4)$$

where

$$0 < t < 1 .$$

The initial temperature is chosen such that it is of the same order of magnitude as the energy difference between the first two configurations. This is done in an attempt to scale the energy difference between successive configurations to something between zero and one. Other choices of temperature schedule are possible and these are currently the topic of research.

## 4.2. Bayesian Analysis

In the case where we have a series of similar *in situ* tests available, but the prediction we wish to make is for a dissimilar test, i.e. one that is far away from all the other tests, or one that has distinctly different flow conditions, we use the Bayesian approach or method of regulariza-

tion to incorporate prior physical information into the model. This approach is based on the fact that quantitative information, such as the range of possible responses, might be available even if there is no actual measurement. Using this information, we can constrain the model to give a response within this range. In this case, we might have some general information about the area, such as a range of a certain response should fall in. This information can be used in the annealing process, by adding a step which makes it more likely that configurations which have a calculated response value in this range will be accepted. The values predicted by the final models may be used to update your estimate for this response.

For example, suppose we want to predict the flow to a borehole in the model at a place far from the present boreholes. We have as prior information that steady state drawdown values have been observed to be between 10 and 20. That is

$$h \in [10,20]$$

We then anneal using the energy function based on the known borehole responses. However, every time we change the configuration we do the following:

1. Add the new borehole to the configuration and calculate the drawdown,  $h$  at that borehole.
2. Compute a new energy function  $Q'$  as follows:

$$Q'(C) = \begin{cases} 0 & \text{if steady state value} = h \in [10,20] \\ [h-10]^2 & \text{if } h < 10 \\ [h-20]^2 & \text{if } h > 20. \end{cases} \quad (4.5)$$

3. Add  $Q'$  to the energy calculated for the known responses,  $Q$ .
4. If  $Q(C') + Q'(C')$  is an improvement over  $Q(C) + Q'(C)$ , keep  $C'$ .  
If not, use usual annealing probability to keep or reject  $C'$ .
5. Remove new borehole and continue.

### 4.3. The Annealing Computer Algorithm

The finite element code, TRINET, simulates steady or transient flow in a fracture network. The code is used to solve for the head distribution as a function of time in the fracture mesh created by the mesh generator in response to a simulated well test. For example, the head measurements from transient constant flux well tests could be simulated with TRINET. Now, the head versus time curves from the simulated well tests are examined and the values of transmissivity and storativity assigned to the elements are scaled in order to achieve as good a match as possible to the observed draw down curves. This corresponds to shifting the well test data to match the Theis type curve (Freeze and Cherry, 1979). We then define the "energy,"  $Q(C)$ , of our current model configuration as the squared sum of the differences between the observed and calculated head values at different locations ( $j$ ) and specified times ( $i$ ):

$$Q(C) = \sum_m \sum_i [H_{obs}(i,j) - H_{sim}(i,j,C)]^2 \quad (4.6)$$

In the case presented here, we do not have a transient test, only a single steady state example. So the energy function is just computed at the points where head is monitored:

$$Q(C) = \sum_j [H_{obs}(j) - H_{sim}(j,C)]^2 \quad (4.7)$$

The computer algorithm can be written in 4 steps.

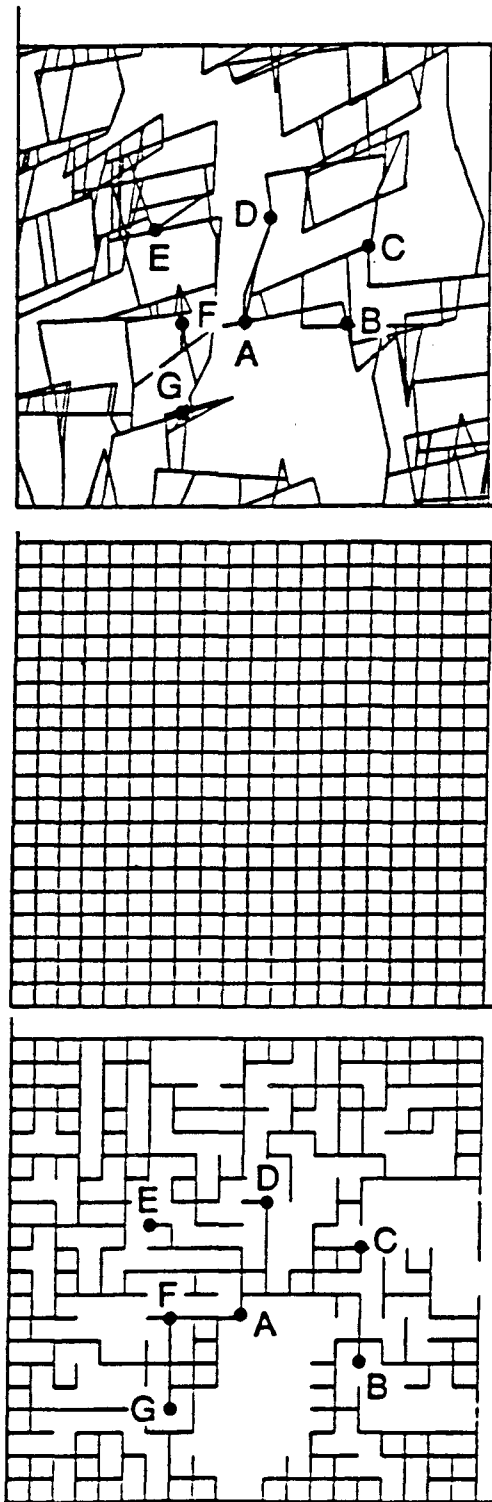
1. Randomly select a beginning configuration from the set  $\Omega$ . Calculate  $Q(C)$ .
2. Randomly select a pipe from  $M$ . If the pipe is in  $C$ , delete it, if the pipe is not in  $C$ , add it. This is the new configuration,  $C'$ .
3. Calculate  $Q(C')$ .  
 If  $Q(C) > Q(C')$ ,  $C \rightarrow C'$ .  
 If  $Q(C) < Q(C')$ , use a random number and the transition probability to decide if  $C \rightarrow C'$  or  $C \rightarrow C$ . If  $Q(C')$  is accepted then  $Q(C')$  becomes  $Q(C)$  for the next iteration.
4. If the end of step  $k$  of the annealing schedule has not been reached, return to 2 and con-

tinue. If the last iteration of step  $k$  was reached, reduce  $T_k$  to  $T_{k+1}$ , return to 2 and continue. If the last step  $k$  of the annealing schedule was reached, stop if no change was made in the configuration after some large number of iterations.

#### 4.4. A Synthetic Example Case

An example of annealing results is shown in Figure 4.1. Here we have created a synthetic fracture system (Figure 4.1) and used it to create synthetic well test data. Then, we create a regular grid to use as a template (Figure 4.1) and apply annealing to find a configuration which matches the synthetic well test data (Figure 4.1). This example shows that the annealed result roughly reproduces the connection between the wells. Major gaps similar to those in the "real" system, have analogous gaps in the annealed system.

A full suite of synthetic cases is being developed to study the effect of template geometry, temperature schedule, weighting functions, etc. (Davey *et al*, 1990). This suite should give us experience that can be applied to cases where the true geometry is really unknown.



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Figure 4.1. (a) A "real" fracture network where we have generated "real" data on well test in hole A and monitoring in holes B through G; (b) Template model; (c) The pattern of conductors resulting from annealing.

## 5.0. MEASURES OF UNCERTAINTY

After annealing has been used to identify model configurations of conductors which match the observed *in situ* behavior, the models can then be used to make predictions of other flow situations. In this example, we first apply boundary conditions to simulate the well tests used for annealing. When annealing is completed, we shut in these wells in the model, and create new boundary conditions appropriate for predicting some new flow conditions. We now wish to determine how good these predictions are. There are two approaches to this problem: prediction error (or cross validation) or sensitivity and relative likelihood. These are discussed below.

### 5.1. Prediction Error

The first approach, and by far the best, is to use the annealed model to make a series of predictions similar to the one which is required, but for which the quantities predicted have actually been measured. In other words, we ignore some of the data and use the model to predict the data we ignore. Then, for each prediction we can calculate a prediction error, and the root mean square prediction error is a direct measure of how well we can make similar predictions. This is the best way because all of the uncertainties in the model itself are integrated into an error which can actually be measured.

For example, suppose we have 5 wells, and a cross-hole well test from another, sixth well to all 5 wells. This gives an observed response  $H_{obs}^i$ , at each well,  $i$ . We would like to know the prediction error associated with using our annealing model to predict the response at a seventh nearby well. For example, if we were using transient well test data, in the general case we could do the following:

1. Leave the well test curve for well  $i$  out of the energy function.
2. Anneal  $n$  times using the other 4 well test curves until the energy is sufficiently low.



3. For each end configuration,  $C_1^{-i} \cdots C_n^{-i}$ , calculate a predicted well test curve for well  $i$ .

These head versus time curves are  $H_1^{-i} \cdots H_n^{-i}$

4. Calculate the prediction error for well  $i$ :

$$PE^2(i) = \frac{\sum_{j=1}^n (H_j^{-i} - H_{obs}^i)^2}{n}$$

The total prediction error is:

$$PE = \left[ \frac{\sum_{i=1}^5 PE(i)}{5} \right]^{1/2}$$

PE is then an estimate of the error involved in using the annealing model to predict the response of any other well in the vicinity. The more different kinds of responses we can use to calculate the prediction error, the more confidence we will have in the predictions that are made. If there are enough cases to allow us to calculate an empirical distribution of prediction errors, then we can put confidence limits on the prediction.

## 5.2. Sensitivity and Relative Likelihood

Unfortunately, a good estimate of prediction error will require a good sample of *in situ* tests. As discussed before under section 1.3 on validation, this is not always available. When prediction error can not be estimated, we can separate the uncertainty into two components: uncertainty in the conceptual model and uncertainty in the parameters. The uncertainty in the conceptual model can be approached by trying different conceptual models. For example, we can compare the zone model to the grid model. However, there are a very large number of possible conceptual models and we must invoke a "peer review" to decide if we have looked at enough of these.

This leaves uncertainty in the parameters of the model. To approach this problem, we can perform studies which vary the parameters to see what effect they have on the prediction. In our case, we have essentially three kinds of parameters: boundary conditions, the value of the con-

ductance assigned to each conductor, and finally the arrangement of conductors in the model. A study of the sensitivity to boundary conditions can be done by trying different scenarios such as a free surface boundary or adding the drifts to the model. As for the conductances, as long as we use models with constant conductance, any prediction of steady flow rate will be directly proportional to the conductance assigned to the elements, so sensitivity to this parameter is known *a priori*.

Finally, we can examine the uncertainty in the arrangement of the conductors, looking at a series of different arrangements and calculating the energy function for each one. The lower the energy, the more likely is the arrangement. Therefore, we can not say what the probability of each arrangement is, instead we can determine the relative likelihood of any predictions that we make. If the energy function we use in the annealing process is a combination of energy functions for different types of observations, we may rank the final solutions differently for each type of response.

There are several ways which we could choose different configurations of conductors to compare. The simplest is to use a series of configurations defined at the end of an annealing process. These configurations are easily available, but they will probably be very similar to each other. Another way to find different configurations is to perform annealing several times, each time starting with a different initial configuration. For example, we could start with different initial arrangements all with the same percentage of elements present, or we could start with different percentages. For a complete treatment, these exercises should be performed. They are not included here because the available data is not complete enough to warrant this effort.

As these different configurations are developed, one would also want to examine them to see if they are alike geometrically. If models started from different initial cases and all develop the same characteristics, we would want to examine these characteristics in light of the geologic and geophysical evidence. For example, the models may all show lack of connection between two points, or they may all show a preference for strike oriented channels over dip oriented channels.

Mathematically, the relative likelihood is calculated as follows. Assume we have an annealed model with end configurations:

$$C_i \quad i=1 \cdots n$$

For each configuration, we know the the energies

$$Q(C_i) \quad i=1 \cdots n$$

and the final temperature, T. So we can compute the relative likelihoods of the i configurations:

$$e^{\frac{-Q(C_1)}{T}} \cdots e^{\frac{-Q(C_n)}{T}}$$

and rank the estimates from most likely to least likely using the relative likelihoods.

## 6.0. PREDICTION OF INFLOW TO THE D-HOLES

### 6.1. Development of a Synthetic Cross-hole Test

At this stage in the hydraulic investigation of the SCV site, there are no formal, well controlled cross-hole well tests available. In order to gain experience with the annealing technique and produce a preliminary estimate of the flow into the D-holes, a synthetic cross-hole test was produced based on a variety of data available for the SCV site. This data consisted of ad hoc cross-hole tests performed by British Geologic Survey (BGS) and the record of heads in the boreholes.

BGS conducted three ad-hoc cross-hole tests by opening W2 and monitoring sections in N3, N4, and W1. Some zones did not respond while others responded very quickly. The test durations were too short to achieve steady-state conditions. At the same time, the transient data was, at best, qualitative because the tests were conducted while the system was in a recovery period due to other perturbations. Also, the test was started without waiting for the packer inflation effect to diminish or for the zone pressure to build up to the environmental head. Therefore, the data is a complex combination of all three effects. We did not attempt to analyze these effects and did not use the transient data directly in the quantitative annealing procedure. However, qualitatively these tests offer valuable information of the major hydrologic features. In comparison to the single hole tests, the scale of these cross-hole tests is much more representative of the overall size of the SCV site and the large hydrologic features.

Based on the transient results plus the record of hydraulic heads in the holes, a synthetic steady state test was fabricated. The steady flowrate from W2 was extrapolated to estimate the steady-state flow rate (Figure 6.1) of 10 liters/min. The corresponding estimation of steady state head in N3, N4, and W1 was found by extrapolating the head values in those holes during the period when W2 was opened for a prolonged period of time (Figure 6.2). Wherever possible, these heads were assigned to specific zones in the boreholes based on the responses observed

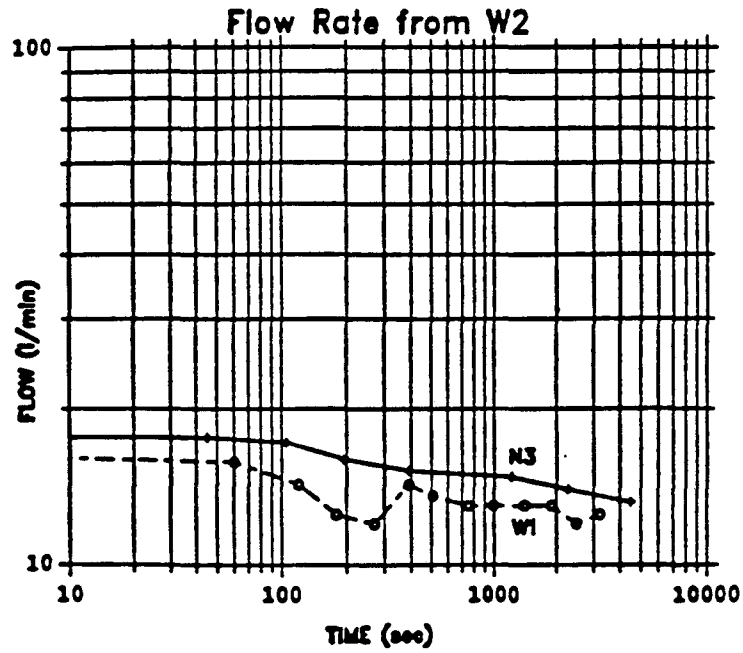


Figure 6.1. Flowrate in W2 recorded during two periods: when N3 was monitored and when W1 was monitored.

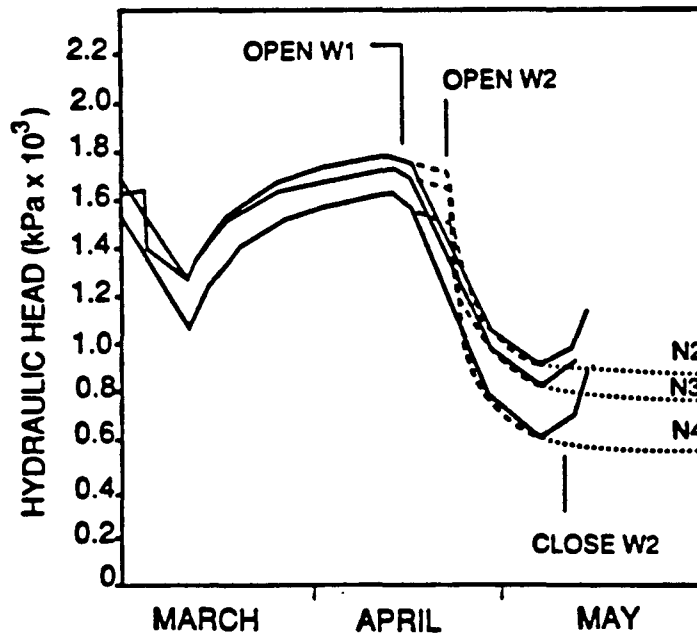


Figure 6.2. Responses in N2, N3, and N4 (entire hole) to opening W2 (from the head records, 1987).

during the ad hoc test. In other words, during the ad hoc tests, the observation holes were divided into several sections. Some sections did not respond to opening W2, whereas others responded very rapidly. In these cases, the responding zone was assigned the observed head.

The annealing case is summarized in Figure 6.3 and in Table 6.0. Figure 6.3 shows the N- and W-boreholes, plus the "fins" that are used to connect the boreholes to the hydrologic zones. Each fin represents the intersection of a zone with the borehole. The black fins are those where we have determined a value of head to use in the calculation of annealing energy based on the ad hoc cross hole tests plus the head record. The gray fins in W2 are held at zero head to simulate the opening of W2. The white fins are those for which we have no record of response to opening W2. In these cases, we could not be sure that there was no response under steady conditions. We only knew that no response was observed under the limited testing period. Therefore, the value of head at these fins is calculated, but not used in the annealing process. When a zone does not respond, the annealing algorithm will disconnect the zone, so uncertainty about lack of response is a significant loss of information.

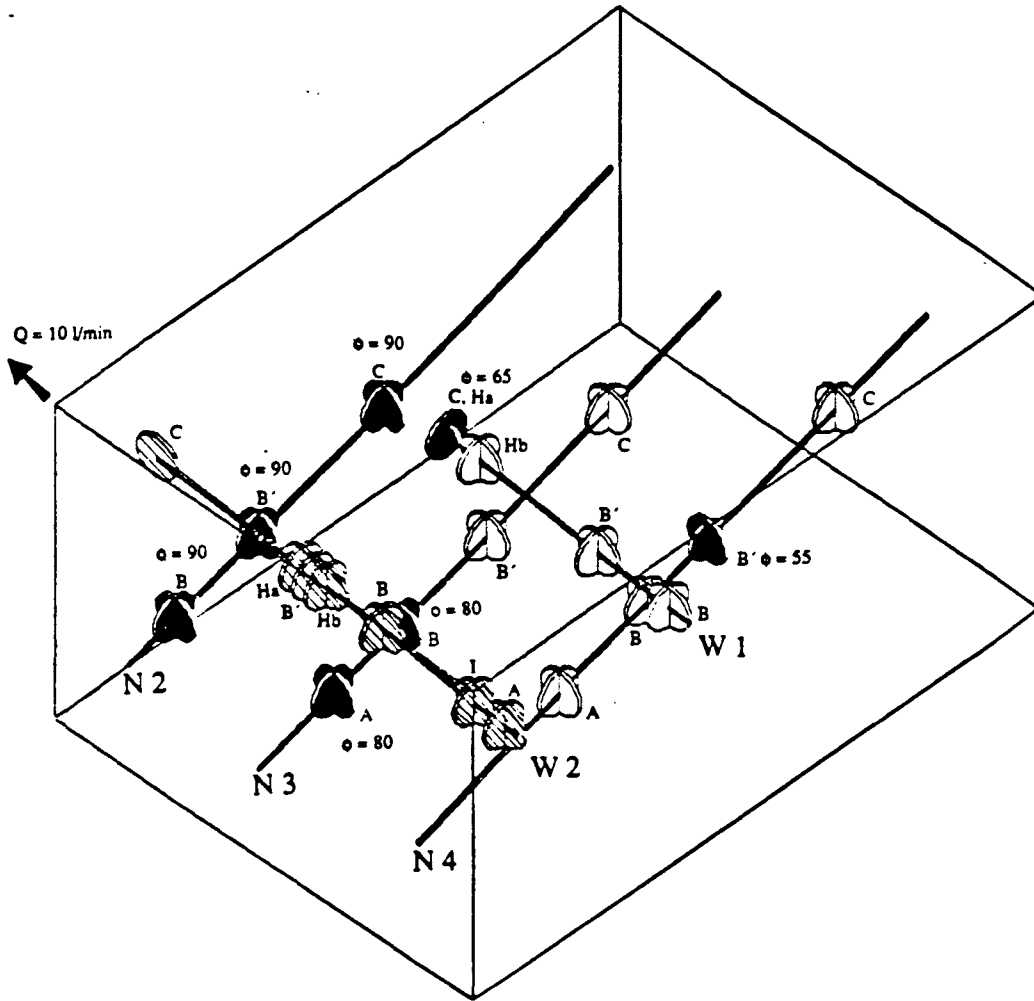
Table 6.1. Annealed Borehole Sections and Head Values




Hole	Depth (m)	Zones	Head (m)	Flow (l/min)
N2	0-207	B,B',C	90	0
N3	131-189	A,B	80	0
N4	87-100	B'	55	0
W1	43-51	H <sub>a</sub> ,C	65	0
W2	0-147	A,B,B',C,H <sub>a</sub> ,H <sub>b</sub> ,I	0	10

## 6.2. Boundary Conditions and the Starting Configuration

There are two possible choices for setting the boundary conditions. A regional mine scale hydrologic model has been constructed (ref. J. Gale in Olsson, 1989). This model could be used to pick out the hydraulic heads at the boundaries of the SCV block and assign these to the zone model. This option was rejected because changes in the properties of the zones model would have a large effect on the boundary heads. Thus, the mine model would have to be directly coupled to the zone model in order to predict the heads on the boundary. This option is not practical

### THE ANNEALING CASE



-  "Observed" heads (m) based on synthetic steady flow from W2, used in calculating the energy for annealing.
-  Value of head not "observed", predicted by annealing.
-  Open hole,  $\phi = 0$

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Figure 6.3. Summary of the synthetic annealing case.

in conjunction with annealing.

An alternative is to base the choice of boundary conditions on head observations in the boreholes. The SCV block is situated in a large zone of depressed heads which represent the steady drainage created by the mine. Therefore, whatever perturbation there are in the SCV block must be superimposed on this existing condition. Shut-in heads throughout and around the SCV block are surprisingly consistent and centered about 200 m. Therefore, we choose the alternative of making the boundary conditions constant head equal to 200 m. Although we do not expect the boundary conditions to be uniform around the edges of the zones, we have too little data to have any more resolution than setting the boundary conditions to one estimated figure. These boundary conditions represent a degree of approximation consistent with the rest of the assumptions we have made so far, and the boundary conditions do not seem to have a large effect on the resolution of conduction patterns in the vicinity of the wells.

Using 200 m head boundary conditions applied directly on the edges of the block and opening W2 to atmospheric pressure, results in the model calculating heads in the other N- and W-holes which are too high compared to the actual heads that were observed when W2 was opened. Further, we know that pressure perturbations in the SCV block are felt much farther away than around the boundary of the SCV block. We conclude that having fixed head boundaries coinciding with the boundary of the SCV block is unrealistic, because these boundaries are hydraulically too close to the N- and W-holes to allow for the observed head drops.

In order to get a better representation of the hydraulic conditions in the block, we connected each zone to boundaries set at 200 m head far away from the block. The connection was made through channels which had conductance equal to that of the channels in the block. The distance of 1000m was chosen so that the simulated heads in N- and W-holes are, on average, close to the observed heads. This distance is larger than the distance to the water table, but using 1000 m is numerically equivalent to setting the permeability of the rock surrounding the SCV block to be much lower than that of the zones. Therefore, we are effectively controlling the head drop to the boundary of the block, but we are not concerned with what actually causes the head



drop outside of the block.

One remaining question is whether to include the drifts themselves in the model in order to incorporate their effect on the heads in the block. We decided against this as a first approximation because the heads in the SCV block very near the drifts are close to 200m. Thus, we would have to surround the drifts with a low permeability zone or they would cause a much larger head drop in the model than is observed. Surrounding the drifts with a low permeability zone would be almost equivalent to not including them, so we left them out.

A modification of the energy function was defined for this case because there is some uncertainty about the boundary conditions. We are not very confident of the the inner and outer boundary conditions because local heterogeneities near W2 will cause the system to behave as if the boundary at W2 is not zero head. If there is a zone of low permeability around the well, the effective head is higher than zero and *vice versa*. We know much less about the outer boundary condition. The relative head structure observed away from these uncertainties is considered to contain more information. So, the inner and outer boundary conditions were shifted up or down such that we could obtain the minimum relative distance to the observed values. In this way, we are putting all the annealing emphasis on the pattern of the head values and not on the absolute value of the heads.

### 6.3. The Energy Versus Iteration Curve

A temperature schedule was chosen as:

$$T_{k+1} = T_k 0.7^k \quad (6.0)$$

with

$$T_0 = 0.1 \quad (6.0)$$

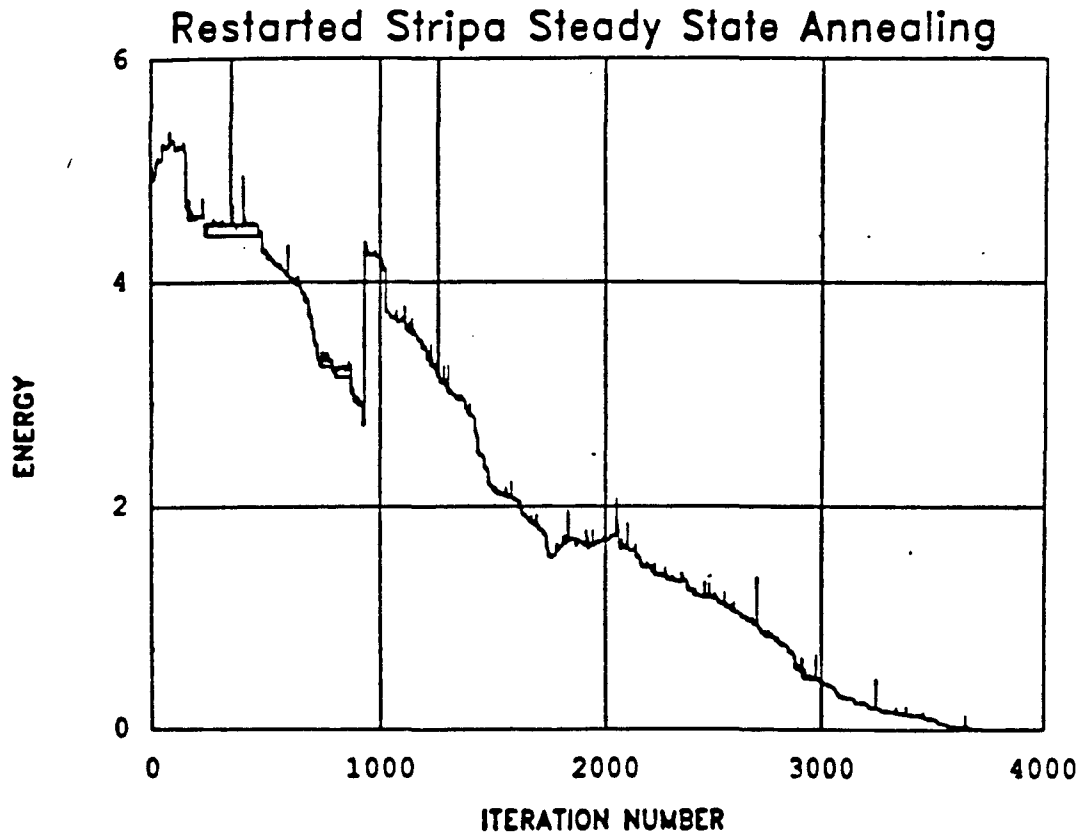
and the temperature was changed every 20 successful iterations. The annealing program was initiated and continued running for 931 iterations during a period of one week. At that time, an inefficient algorithm in the program was detected which was causing the annealing procedure to occasionally retain the previous configuration unnecessarily. This is evident in Figure 6.4

between iteration 1 and 931 by the regions on plot where there are two values of energy plotted for the same iteration. At the same time, we found that the head in N3 had been left out of the energy calculation. Hence, the process was stopped and restarted with a refreshed temperature schedule, the current minimum energy fracture configuration, the error corrected, and the fourth well included. Recall from the discussion of annealing theory that a restart is justified because these configurations are random Markov Fields.

The added well sharply increased the energy level at this point (Figure 6.4, iteration 932). With the process correctly functioning, the procedure continued for another five days until the temperature schedule was exhausted and the procedure terminated normally at iteration number 1813 with energy 1.74. To see if we could get the energy closer to zero, the annealing program was restarted with an extended temperature schedule at the current configuration. Also, the annealing program was altered to increase the number of iterations at each temperature from 20 to 50. This change is reflected in the decreased slope of the receding energy function. The process was stopped at iteration number 3749 and energy 0.005661.

Figures 6.5 gives the final configuration of the channels in each zone. Each figure shows the complete set of possible channels, and the configuration of channels at the last iteration.

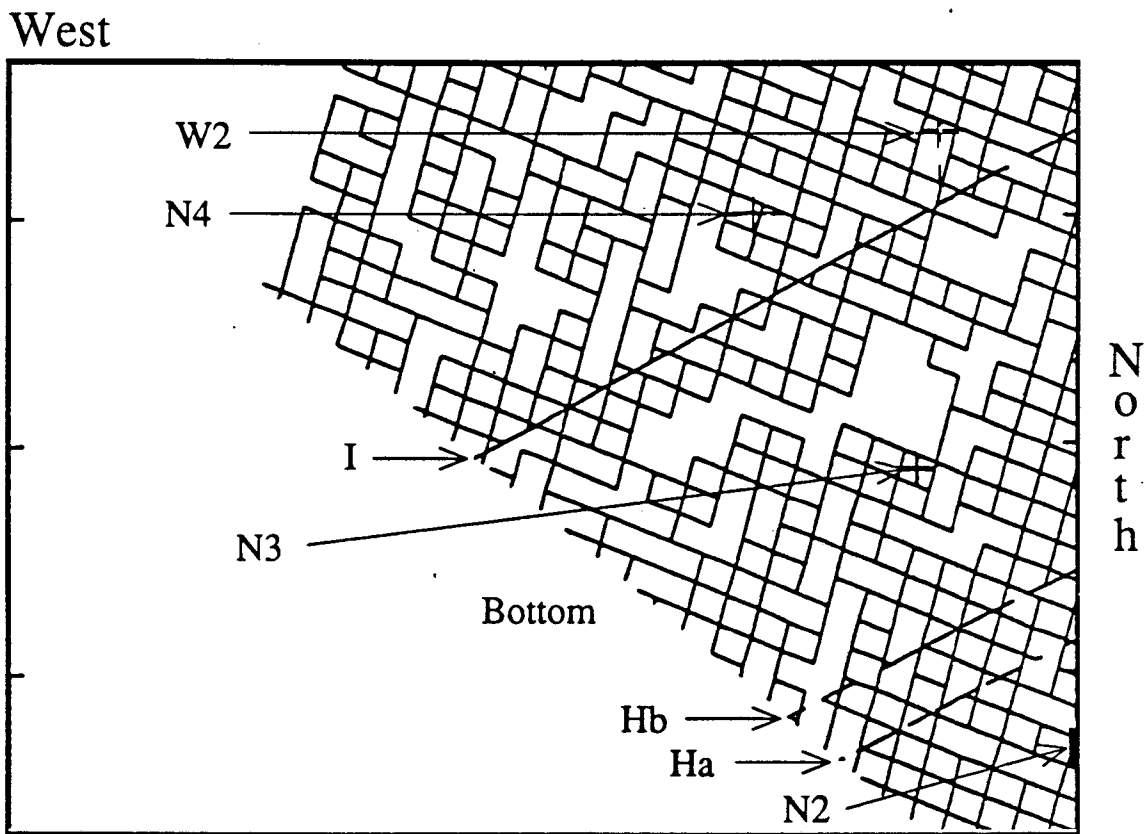
Table 6.2 gives the heads that were calculated with the final configuration resulting from annealing. We see that the annealing routine has managed to match the observed heads very well. The other values of head can be compared to those measured when the D-holes are brought to near zero pressure. However, there is a great deal of uncertainty about the boundary conditions and these predictions are directly related to the applied boundary conditions.



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Figure 6.4. Record of energy versus iteration for the SCV synthetic well test case.

### Zone A



ESD-907-0012

Figure 6.5. Final configuration of grid elements in zone A.

Table 6.2. Annealing results at the final iteration = 3749

Hole	Zone	Observed Heads	Predicted Heads
N2	B,B',C	90	90
N3	A,B	80	79
	B'	none	65
	C	none	65
N4	B'	55	55
	C	none	83
	B	none	49
	A	none	49
W1	H <sub>a</sub> ,C	65	65
	H <sub>b</sub>	none	65
	B'	none	65
	B	none	65

#### 6.4. Calibration to W2 Flow (Shifting)

At the end of the annealing process, we have determined several configurations of conductors within the zones all of which result in matching the observed head data extremely well. The match has been achieved solely by arranging the conductors. As all the channels have the same conductance,  $k_A$ , any value of  $k_A$  will result in the same head distribution. Thus, at this point we must calibrate the model such that it will predict the correct value of flow from W2. To do this, we use the annealed model to calculate the flow from W2,  $QW2_A$  with all conductances equal to  $k_A$ . Then we take the ratio of measured flow to calculated flow to find the conductance of all the channels,  $kW2$  which would produce the correct amount of flow into W2 ( $QW2_M$ ).

$$kW2 = k_A \cdot \frac{QW2_M}{QW2_A} \quad (6.0)$$

For the final configuration of the model, the value of  $QW2_A$  is  $3.14 \text{ m}^3/\text{s}$ , and  $k_A$  is  $0.42 \text{ m}^3/\text{s}$ , so for the case of  $QW2_M = 10 \text{ l/min} = 1.67 \times 10^{-2} \text{ m}^3/\text{s}$ , we require that  $kW2 = 2.2 \times 10^{-5} \text{ m}^3/\text{s}$ .

Now we rearrange the numerical model, closing the hole W2, and adding the open D-holes and calculating the outflow from the D-holes. The resulting calculation of inflow to the D-holes for the final configuration is 8.9 liters/min.

However, by the scaling equation above, the flow into the D-holes is directly proportional to the flow from W2. This indicates that the prediction of D-hole inflow is extremely sensitive to the measurement of flows. In a good annealing case, we would have been able to use the transient curves or flow rate data in the calculation of Energy. In this case, the calculation of flow depends on a single measurement of flow, and we suspect that this measurement is anomalous in that the transmissivity in W2 is much higher than the other holes. This points out that other flow data available for the SCV block would be very useful in modifying this prediction and this is discussed below. Later work (Davey *et al.*, 1990) has confirmed the utility of cross-hole data in determining fracture pattern.

### 6.5. Modification of the Prediction Using Other Inflows

One further set of data is available to aid in the prediction of inflow to the D-holes. This data was also collected on an ad hoc basis by BGS (D. Holmes, personal communication), and consists of measurements of outflows from the other N- and W-holes after they had been left open for periods of time. Table 6.3 gives this data.

Table 6.3. Open Hole Inflows to N- and W-holes

Hole	Flow (l/min)	Comment
N2	0.60	Measured flow after 6 hours open
N3	0.45	Measured flow after 4 hours open
N4	2.55	Measured flow after 2 hours open
W1	1.30	Measured flow after 2 days of chemical sampling
W2	12.0	Measured flow after 3 hours open

First, we note in Table 6.3 that the measured inflow to W2 is 12 l/min. Our best estimate of the steady flow to W2 is 10 l/min, or about 83% of that in Table 6.3. This is because the values in Table 6.3 do not yet reflect steady flow. To correct for this, each of the measurements of flow is reduced by 17%. We call these flows  $Q_{iM}$  where i stands for the holes, N2, N3, N4, W1, and W2.

Now, we can use the final annealed configuration of channels with conductance,  $k_A$ , to calculate the inflow into each of these holes,  $Q_{iA}$ . To do this we simply close W2 by making the nodes at W2 internal nodes, then sequentially open each of the other holes by assigning their nodes zero head. In each case we calibrate the channel conductance in the same manner as previously described such that the model correctly predicts  $Q_{iM}$ :

$$k_i = \frac{Q_{iM}}{Q_{iA}} k_A \quad (6.0)$$

This results in five different predictions of channel conductance, which in turn results in five different predictions of D-hole inflow. These predictions are given below in Table 6.4.

In summary, our prediction of inflow to the D-holes has mean 3.15 l/min and a coefficient of variation (standard deviation divided by the mean) equal to about one. To reinforce the point that the resulting prediction is dominated by the flow measurements, we note that the mean and standard deviation of the predicted inflow is close to the mean and standard deviation of the adjusted observed flows from the five holes. We do not know the form of the inflow distribution function, but if we make a guess that inflows are distributed log-normally, then we can calculate that the flow will be between 0.44 and 11.4 with 95% confidence.

Table 6.4. Predictions of D-Hole Inflow Based on Annealing and Measured N- and W-Hole Inflows.

Hole	Measured Flow (l/min)	Adjusted Flow (l/min)	$k_i/k_A$ $\left[ \frac{l/m}{m^3/S} \right]$	Predicted D-hole inflow (l/min)
N2	0.60	0.50	0.16	1.3
N3	0.45	0.37	0.12	0.6
N4	2.6	2.1	0.67	3.4
W1	1.3	1.1	0.35	1.3
W2	12.0	10.0	3.2	8.9
Mean	3.4	2.8	.89	3.1
Standard Deviation	4.9	4.1	1.3	3.1

### 6.6. Estimation of Prediction Error

Recall that to calculate prediction error, we need more than one test, say  $N$  tests. These  $N$  tests would each consist of a measured flow rate plus measured head responses. Then we could anneal the model  $N$  times, each time leaving out the data from one of the tests. With each model we would predict the flow for the test that was left out, calculate the difference between the observed flow and the predicted flow, and take the root mean square of these differences to get the estimate of prediction error.

In our case, we only have draw-down data from one test, but we have five different measurements of inflow. Hence, to make an estimate of prediction error, we divide the into two types, head and flow data. We anneal with the head data alone to get a channel configuration. Then we develop five different models by calculating channel conductance with only four of the inflows at a time. Thus, we average the conductances,  $k_i$ , from Table 6.4 leaving out one value (the  $i$ th value) at a time to get  $\langle k_{i0} \rangle$ . Then, we use this value to compute the flow into the hole we left out of the average. The root mean square of the differences between predicted and observed flux for each case is the estimate of prediction error. These values are given below in Table 6.5.

Table 6.5. Prediction Error

Hole Left Out	$\langle k_{i0} \rangle$	Predicted $Q_i$ (l/min)	Observed $Q_i$ (l/min)	Error (l/min)
N2	1.1	3.4	.5	2.9
N3	1.1	3.4	.37	3.0
N4	.96	3.0	2.1	.90
W1	1.0	3.3	1.1	2.2
W2	.33	1.0	10.	9.
Estimated Prediction Error				4.6



## 7.0 SUMMARY OF RESULTS AND CONCLUSIONS

Based on the preliminary data available at this time, the inflow predicted for the D-holes is 3.15 l/m with a prediction error equal to 4.6 l/m. The coefficient of variation for the individual flows to the D-holes is about 1.0. The results are most sensitive to the measurements of inflow. We think that this is particularly due to the fact that the annealing case was so limited and that only steady state data was used. Based on synthetic examples, we expect that multiple transient annealing will be able to discern channel patterns much more effectively than steady state annealing. However, because the estimate is sensitive to flow measurements and it is the flow we are predicting, we have confidence that this approach gives answers that make more sense.

We used an energy function defined with the relative distance to the observed head values, that is we shifted the boundary conditions to minimize the energy. It is of interest to compare these results to the case where the absolute value of head difference is used instead. However, at the last iteration the shift was small (about 1.4 m) so this change in approach will probably not have a large effect on the result. Even more enticing is the idea of using the configuration at each iteration to sequentially model flow into each of the N- and W-holes. In this case, we could define an energy function which includes terms for the differences between predicted and measured flow, as well as head. Thus, at each iteration we would run the flow code five times instead of only once. A weighting function would probably be needed in order to account for the fact that the units and magnitude of head differences are different than the units and magnitude of flux differences. In this case, we would expect annealing to be more sensitive to the heterogeneities near the wells which control the inflows, so we would not have had to use the boundary condition shifting. Although this approach is more CPU intensive, it may converge faster and would be worth trying.

Questions remain about weighting some data over others and the effect of different starting configurations of the model. Davey *et al.* (1990) reports on such studies and shows that the initial

percentage of conductors has little effect on the final percentage. Also, it is not surprising these synthetic cases show much better resolution of pattern when the template reproduces the orientations of the real conductors. This fact supports the need for the *a priori* definition of the template using geology and geomechanics approaches.

Finally, these results indicate that we may have over-discretized the channel network of the template. Possibly one of the reasons that the annealing did not result in any pattern definition is that the network simply allowed too many degrees of freedom. There is a need to look at discretization versus information content in the well tests. This also could be done with synthetic examples.

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