

## Scaling issues in snow hydrology

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### Abstract:

The concept of scale can be used to quantify characteristic lengths of (a) a natural process (such as the correlation length of the spatial snow water equivalent (SWE) variability); (b) a measurement (such as the size of a snow density sample or the footprint of a satellite sensor), and (c) a model (such as the grid size of a distributed snow model). The different types of scales are denoted as process scale, measurement scale and model scale, respectively. Interpolations, extrapolations, aggregations, and disaggregations are viewed as a change in model scale and/or measurement scale.

In a first step we examine, in a linear stochastic analysis, the effect of measurement scale and model scale on the data and the model predictions. It is shown that the ratio of the measurement scale and the process scale, and the ratio of the model scale and the process scale are the driving parameters for the scale effects. These scale effects generally cause biases in the variances and spatial correlation lengths of satellite images, field measurements, and simulation results of snow models. It is shown, by example, how these biases can be identified and corrected by regularization methods. At the core of these analyses is the variogram. For the case of snow cover patterns, it is shown that it may be difficult to infer the true snow cover variability from the variograms, particularly when they span many orders of magnitude.

In a second step we examine distributed snow models which are a non-linear deterministic approach to changing the scale. Unlike in the linear case, in these models a change of scale may also bias the mean over a catchment of snow-related variables such as SWE. There are a number of fundamental scaling issues with distributed models which include subgrid variability, the question of an optimum element size, and parameter identifiability. We give methods for estimating subgrid variability. We suggest that, in general, an optimum element size may not exist and that the model element scale may in practice be dictated by data availability and the required resolution of the predictions. The scale effects in distributed non-linear models can be related to the linear stochastic case which allows us to generalize the applicability of regularization methods. While most of the paper focuses on physical snow processes, similar conclusions apply and similar methods are applicable to chemical and biological processes. Copyright © 1999 John Wiley & Sons, Ltd.

**KEY WORDS** scale; scaling; aggregation; snow cover patterns; regularization; distributed snow models; fractals; effective parameters; representative elementary area; subgrid variability; snow water equivalent

### INTRODUCTION

This paper addresses two questions: (1) How can we measure and represent snow processes at different scales? and (2) How can we aggregate and disaggregate spatial snow data?; These are very broad questions indeed and have ramifications in three main areas of snow hydrology: (i) In the quest for revealing the true nature of snow processes, more specific questions include: What is the nature of spatial snow variability?

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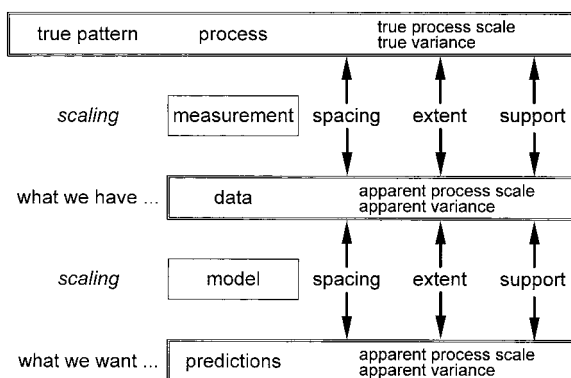


Figure 1. Scale effects in the measurements cause biases in the data; scale effects in the models cause biases in the model predictions. Scaling or change of scale is depicted as double arrows

and: How does it change with scale? (ii) In the context of measurements and data collection more specific questions include: At which scale should we measure?; How can we interpret data measured at a given scale?; and How can we best design a snow measurement network? (iii) In modelling, there are issues related to spatial estimation in general. For example: How can we interpolate/extrapolate from measurements?; How can we aggregate/disaggregate measurements?; or more generally, How can local observations be transferred to larger scales? In modelling, there are also issues related to distributed physically based snow models, such as How can we represent spatial processes in these models; How can we represent subgrid variability?; and What is the optimum model resolution? Clearly, there are more questions here than are likely to be answered in the near future. This paper attempts to contribute to a more coherent understanding of these issues by proposing a framework within which we can deal with these issues.

On closer examination, and from a more general perspective, most of these issues arise because the *scale* at which the data are collected is different from the scale at which the predictions are needed. Figure 1 shows a sketch of how natural snow variability, data and model predictions are related in general. At the top of Figure 1 is the true spatial variability, i.e. the true snow hydrological processes which, however, will never be known in full detail. These natural processes possess certain properties such as spatial patterns of a variable, a true variance, a true correlation length, etc. To obtain information on these processes, measurements are made, either by data collection of a standard hydrologic network, in research catchments, at study plots or by remote sensing techniques. The measurements produce data. The important thing is that the data will not accurately portray the natural variability because of a number of factors. Among these factors are instrument error (which will not be dealt with in this paper) and the spatial dimensions of the instruments (which is the focus of this paper). The patterns of the data will be different from the true patterns and so will their statistical moments. For example, the variance estimated from the data (i.e. the apparent variance) will, in general, be different from the true variance. In a second step, the data are combined by some sort of model to produce predictions. Again, the predictions will in general be different from the data due to a number of factors related to the properties of the model, such as the spatial dimensions of the model or model elements. The apparent variance of the predictions will, in general, be different from the apparent variance of the data. This situation is a very general framework which is probably applicable to any branch of scientific enquiry. Here, we will use this framework to quantify the biases and effects of the various space scales involved in snow hydrologic research.

It may be useful to start with a definition of scale and scaling. We will adopt the definitions proposed by Blöschl and Sivapalan (1995) and Blöschl (1999). The term 'scale' refers to a characteristic length or time. Often, it is used in a qualitative way as in 'a small scale phenomenon' but here we will use it quantitatively to refer to a space dimension. We will propose a 'process scale' that relates to a spatial dimension of natural variability (such as the spatial variability of snow water equivalent); a 'measurement scale' that relates to a

spatial dimension of an instrument or a measurement structure (such as a snow lysimeter); and a 'model scale' that relates to a spatial dimension of a model in a general sense (such as a distributed snow model). These definitions are consistent with the sketch in Figure 1.

We will use the term 'scaling', to denote a 'change in scale'. In Figure 1, there are two changes in scale, from the true process to the data (i.e. scaling by the measurement); and from the data to the predictions (i.e. scaling by the model). The purpose of this paper is to quantify the effects of scaling due to the measurements and the effects of scaling due to the model. Aggregation, disaggregation, extrapolation and interpolation always imply a change of scale and hence we will subsume these things under the term scaling. It should also be noted that the usage of 'scaling' adopted here is one of two usages that can be found in the literature. The other usage of scaling relates to some sort of similarity or scale invariance such as in 'scaling snowdrift development rate' (Lever and Haehnel, 1995), 'scaling Richards equation' (Kutilek *et al.*, 1991) or 'scaling in river networks' (Foufoula-Georgiou and Sapozhnikov, 1998). Although we will also touch on the issue of scale invariance in this paper, for clarity, we will strictly use scaling to denote a change of scale only.

There are a number of factors that complicate scaling in snow hydrology. First of all, the extreme spatial variability of the hydrologic environment greatly complicates the spatial estimation problem (see e.g. Elder *et al.*, 1989). If a snow pack is examined in great detail, the number of possible flow paths for melt water or gas exchange becomes enormous. Along any path the shape, slope and boundary roughness may be changing continuously from place to place and these factors also vary in time as the snow becomes wet. Similar complexity occurs at the catchment and the regional scales. Because of these complications, it is not possible to describe some hydrologic processes with exact physical laws. The scale at which a conceptualization is formulated then becomes critically important and the transfer of information across scales becomes very difficult. Second, when seeking a quantitative description of the natural system one must either adopt a deterministic or a stochastic framework (or a combination thereof). If the spatial patterns of the hydrologic process are 'consistent' enough (i.e. exhibit 'organization') a deterministic analysis is warranted while for a large degree of disorder (i.e. 'randomness') there is usually merit in probabilistic treatment. Unfortunately, hydrologic processes show elements of both chance and structure which complicates both types of analyses (Blöschl *et al.*, 1993). Closely related to organization is the non-linear behaviour of snow processes that complicates scale issues. Third, in many instances there is an enormous contrast in scale between the available data and the prediction required, which may exceed several orders of magnitude. This large contrast may imply that different processes become operative at different scales, and empirical expressions derived for the smaller scale no longer hold true at the larger scale (see e.g. Seyfried and Wilcox, 1995). Clearly, one would not expect processes that control crystal growth at the millimetre scale to be also important at the regional scale where climatic effects are probably more relevant.

There have been a number of reviews of scale issues in hydrology in general (e.g. Blöschl and Sivapalan, 1995) and a number of collection of papers of conference proceedings on the same topic (e.g. Rodríguez-Iturbe and Gupta, 1983; Gupta *et al.*, 1986; Kalma and Sivapalan, 1995; Blöschl *et al.*, 1997). Here, we will build on this work with a focus on snow hydrology.

## THE CONCEPT OF SCALE

### *Process scale*

As mentioned above the 'process scale' relates to a spatial dimension of the natural variability of some snow-related variable. While in principle, there are many possible ways of defining a process scale, it is most frequently defined as the correlation length (or some related measure). The correlation length is based on the variogram  $\gamma$ , which is usually estimated from the experimental variogram

$$\hat{\gamma}(h) = \frac{1}{2n} \sum \{ [Z(x) - Z(x+h)]^2 \} \quad (1)$$

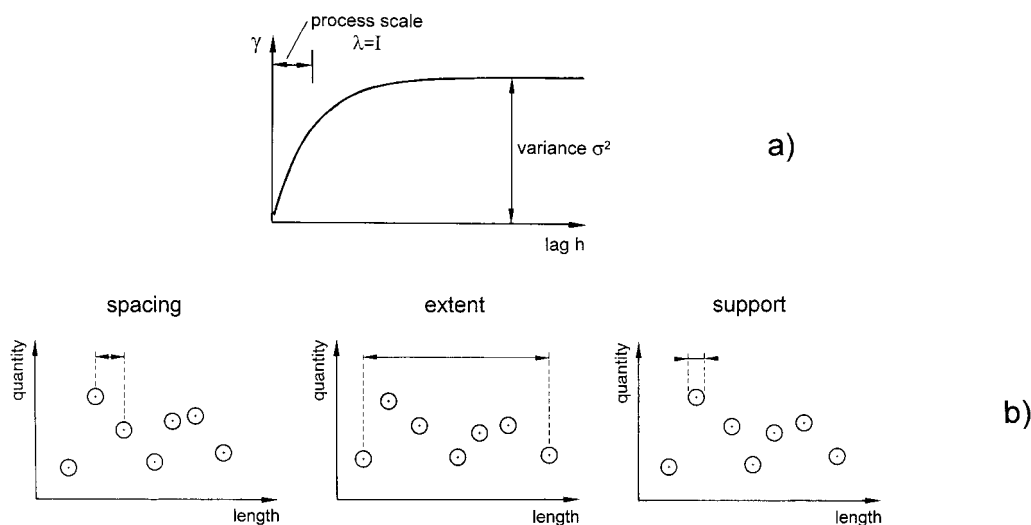


Figure 2. (a) Definition of the process scale (correlation length,  $\lambda$ , or integral scale,  $I$ ). (b) Definition of the measurement scale and the model scale. The scale triplet (spacing, extent and support) can apply to measurements and to models. From Blöschl and Sivapalan (1995)

where  $Z$  are the data (such as snow water equivalent (SWE) or snow depth data) at locations  $x$  and  $x + h$ , and  $h$  is the distance or lag between two data points. In equation (1), the lag  $h$  is subdivided into classes, the summation refers to an individual class and  $n$  is the number of data pairs for each class. The experimental variogram is usually fitted by some analytical function (in the processes of inferring the population statistics from the sample statistics) such as an exponential function of the form

$$\gamma(h) = \sigma^2(1 - \exp(-h/\lambda)) \quad (2)$$

where  $\sigma^2$  is the (spatial) variance of  $Z$  and  $\lambda$  is the correlation length (Figure 2a). In some instances, the integral scale  $I$  is used instead of the correlation length, which is defined as

$$I = \int_0^{\infty} 1 - \frac{\gamma(x)}{\sigma^2} dx \quad (3)$$

For an exponential variogram (equation (2)) the integral scale is equal to the correlation length. For different types of variograms, the integral scale may be a more objective measure of the process scale than the correlation length.

The correlation length (or integral scale) can, strictly speaking, only be derived for a stationary process. For a stationary process, the variogram flattens out at large lags  $h$  (as shown in Figure 2a), which implies that a variance exists. For a non-stationary process, the variogram keeps increasing with large scales. Hence, the variance approaches infinity and the correlation length (or integral scale) also approaches infinity or does not exist. However, it is sometimes possible to assume local stationarity (i.e. the variogram flattens out at some lag after which it increases) as a working hypothesis, in which case a correlation length (or integral scale) can be estimated.

The correlation length (or integral scale) is a measure of the average distance over which the variable is correlated. Equation (2) suggests that for distances shorter than the correlation length the correlations are better than  $r^2 = 0.37$  where  $r^2$  is the coefficient of determination. Indeed, the process scale (correlation length) captures the scale of the continuity of a natural variable such as SWE. If the variable is very continuous, i.e. varies smoothly in space, the process scale is large. If the variable is discontinuous, i.e. with rapid fluctuations over short distances, the process scale is small. A smoothly varying variable will exhibit

large uniform patches in space while a rapidly fluctuating variable will exhibit small uniform patches. For example, the process scale of snow covered area (a binary variable snow covered/snow free) is directly proportional to the average size of the snow free and snow covered patches. More details on the variogram and on correlation scales can be found in the extensive literature on geostatistics (e.g. Journel and Huijbregts, 1978; Isaaks and Srivastava, 1989).

### Measurement scale

Following Blöschl and Sivapalan (1995) we suggest that the measurement scale consists of a scale triplet: spacing, extent, and support (Figure 2b). ‘Spacing’ refers to the distance between samples; ‘extent’ refers to the overall coverage of the data; and ‘support’ refers to the integration volume or area of the samples. All three components of the scale triplet are needed to uniquely specify the space dimensions of a measurement or instrumental setup. For example, for a snow course the scale triplet may have typical values of, say, 100 m spacing (between the samples), 1 km extent (i.e. the length of the course), and 10 cm support (the diameter of the snow density probe). Similarly, for a remotely sensed image, the scale triplet may have typical values of, say, 30 m spacing (i.e. the pixel size), 10 km extent (i.e. the overall size of the image), and 20 m support (i.e. the ‘footprint’ of the sensor). The footprint of the sensor is the area over which it integrates the information to record one pixel value. It is usually on the order of the pixel size but not necessarily identical to it.

In the two-dimensional case it is sometimes useful to relate the spacing, the extent, and the support to areas which is a logical extension from the one-dimensional case in Figure 2b:

$$a_{Spac} = \sqrt{A_{region}/n} \quad (4)$$

$$a_{Ext} = \sqrt{A_{region}} \quad (5)$$

$$a_{Supp} = \sqrt{A_{aggreg}} \quad (6)$$

where  $a_{Spac}$ ,  $a_{Ext}$ , and  $a_{Supp}$  are the spacing, the extent, and the support respectively.  $A_{region}$  is the overall size of the region of interest,  $n$  is the number of samples in this region, and  $A_{aggreg}$  is the area over which a sample aggregates (or integrates). This paper will focus on the two dimensional case.

### Model scale

Again following Blöschl and Sivapalan (1995) we suggest that the model scale consists of a scale triplet (spacing, extent, and support, Figure 2b) very similar to that of the measurement scale. The difference is that the model scale is related to the spatial properties of the model (in a general sense) rather than those of the measurements. For example, for a spatially distributed snow model the scale triplet may have typical values of, say, 25 m spacing (i.e. the grid size), 1 km extent (i.e. the size of the catchment to be modelled), and 25 m support (the grid size). The support is the spatial dimension over which the variables in each grid cell are representative which is equal to the grid size in most distributed models. In regional mapping (which is a model in a general sense) of snow depths the scale triplet may have typical values of, say, 10 m spacing (i.e. the resolution to which the map is produced), 10 km extent (i.e. the overall size of the map), and 10 cm support (which is the area which each point in the map is representative of). In this example, the map shows point values of snow depths (rather than averaged values), hence the support is very small. The support in maps is sometimes termed ‘grain’. Clearly, it is possible to draw two maps of, say, snow depth of the same area, one map showing point values (very small support) and the other map showing values averaged over a certain area (i.e. larger support). The latter map will be smoother than the former.

### Scaling

Here we are interested in the change of the scale, i.e. how will the measurement spacing, extent and support change the true pattern to be reflected in the data; and how will the model spacing, extent and support change the data to be reflected in the predictions (Figure 1). The basic idea of Blöschl and Sivapalan (1995) and Blöschl (1999) is that there is some similarity between these two steps. Generally, if the spacing of the data is too large, the small scale variability will not be captured. If the extent of the data is too small, the large scale variability will not be captured and will translate into a trend in the data. If the support is too large, most of the variability will be smoothed out and the data will appear very smooth. It is clear that some sort of filtering is involved, i.e. the true patterns are filtered by the properties of the measurement which is reflected in the data. It is also clear for dimensional reasons that the effect of the filter will be closely related to the *ratio* of the measurement scale and the process scale. So, more strictly speaking, we should say that 'If the spacing of the data is too large as compared to the process scale, the small scale variability will not be captured'. Analogous statements apply to the extent and the support. The model scale has similar effects. If the spacing of the model elements is too large as compared to the process scale, most of the variability will appear as noise in the predictions. Analogous results apply to the extent and the support of the model. This change of scale, conceptualized as a filter, has been quantitatively discussed in Cushman (1984, 1987), Beckie (1996), Federico and Neuman (1997) and Blöschl (1999), and a more general discussion is provided in Blöschl and Sivapalan (1995).

There are a number of ways of conceptualizing the filters mentioned above. These can be linear filters or non-linear filters; and the analysis can be moulded in a stochastic framework or in a deterministic framework. In snow hydrology, out of the four possible combinations, the two most important ones are a linear stochastic analysis (i.e. mainly geostatistics) and a non-linear deterministic analysis (i.e. mainly distributed physically-based modelling). We will discuss these two most important cases in the following sections.

## LINEAR STOCHASTIC ANALYSIS

### Framework

There are two fundamental assumptions on which this type of analysis rests:

First, the process is linear, i.e. it aggregates linearly or, in other words, simple arithmetic averaging applies:

$$E(f(x)) = f(E(x)) \quad (7)$$

where  $E$  is the mathematical expectation,  $x$  is location and  $f$  is a function or a variable. In snow hydrology there are many processes that do average linearly and many other processes that do not. If  $f$  is a variable, linear averaging is applicable if a conservation law (of mass or energy) holds. For example, if we measure SWE at a 1 m grid in a catchment and calculate the total SWE volume in the catchment from this and, in a second step, aggregate the measurements to a 10 m grid and, again calculate that total SWE in the catchment, these two methods will give the same results due to conservation of mass. In this example,  $x$  in equation (7) relates to the 1 m grid coordinates,  $E(x)$  to the 10 m grid coordinates,  $f(x)$  is SWE at the 1 m grid,  $f(E(x))$  is SWE at the 10 m grid, and  $E(f(x))$  is SWE at the 1 m grid averaged over  $10 \times 10$  m. However, if we aggregate snow albedo to different grid sizes in a physically realistic manner, the average values over the catchment of the two methods will not necessarily be the same. This is because snow albedo is not conservative. If  $f$  is a function representing a measurement set up, there are instruments that do average linearly (such as a raingauge), there are instruments that average linearly in good approximation (such as TDR or satellite sensors), and there are instruments that aggregate non-linearly. If  $f$  is a function in a model representing a hydrologic process (such as Darcy's law), then unfortunately equation (7) often does not apply as most hydrologic processes do not average linearly. In Darcy's law, for example, the average hydraulic conductivity over an area does not give the average flux over the same area, i.e. equation (7) does

not hold. In such cases one may resort to non-linear stochastic analyses (e.g. Gelhar, 1993) or to non-linear deterministic models (see later in this paper).

The second assumption is that the variable to be examined is a second order stationary random variable. Random means that any sort of organization that may be present in the spatial patterns may be neglected. It is clear that catchments are highly organized and, often, snow processes show (organized) linear features. However, in a first approximation it may be useful to assume that snow processes are random in space and to neglect other controls. The assumption of second order stationarity essentially implies that the variogram (equation (1)) does not change with location in the catchment or in the region. It does not necessarily imply stationarity (in the mean), i.e. there is no need that the variogram flattens out at some lag. Second order stationarity may be a reasonable assumption in many applications. However, in some applications, such as network planning, the assumption of second order stationarity does not make sense. In a stochastic approach to network planning the idea is usually that regions with small scale variability (small process scale) are densely sampled (small spacing) while regions with large scale variability are less densely sampled. Clearly, this is inconsistent with the assumption that the variogram (and hence the process scale) does not change with location.

Based on these two assumptions we can now apply standard geostatistics (e.g. Journel and Huijbregts, 1978) or, equivalently, linear filtering theory for random fields (Wiener, 1966) to obtain a filtered random field, i.e. a random field obtained from aggregating, disaggregating, extrapolating or interpolating data points. As mentioned above, aggregation, disaggregation, extrapolation and interpolation (i.e. a change of scale by the measurement or the model) can be viewed as filtering. We will not go into the details of this theory here, which are contained in a rich literature on the subject (see e.g. Journel and Huijbregts, 1978; Vanmarcke, 1983; Isaaks and Srivastava, 1989; Journel, 1993; Armstrong, 1998). We will only present some of the results that are most relevant to snow hydrology in an intuitive way. There are three main results one would be interested in and that can be obtained from geostatistics: (a) the most likely spatial pattern of the variable of interest, (b) spatial patterns with the most realistic variability, and (c) the moments of the patterns. For (a) there are a wide range of geostatistical interpolation methods (kriging and varieties thereof) that are also able to handle the effect of support (e.g. block kriging). These are best linear unbiased estimators that, however, have the disadvantage of tending to smooth out the patterns, i.e. the interpolated patterns are smoother than the real ones. There is a class of interpolators that do not smooth the patterns that are termed stochastic simulations (b). However, this is at the expense of not producing the best interpolations. Stochastic simulations (such as sequential indicator simulation and others) produce a number of equally likely realizations of the patterns, each of them having the same spatial variability (i.e. variogram) as the true pattern. There are software packages available for both classes of methods (e.g. GSLIB, Deutsch and Journel, 1997; SURFER, Golden Software, 1998). In this paper, for clarity, we will focus on (c), i.e. the statistical moments, and how they change with scale. The statistical moments we examine are the mean,  $E(\cdot)$ , the spatial variance,  $\sigma^2$ , and the correlation length,  $\lambda$ . Specifically, we will examine which *biases* will be introduced by a change of scale but we will not examine the random errors (estimation variance or reliability) introduced by a change of scale.

### *Regularization*

There are a range of methods for estimating the filtered moments. These include (a) the spectral method (where the random function is multiplied with the filter function in the frequency domain); (b) filtering in the space domain (simply by running a moving average filter over the data, and estimating the moments from the filtered data); and (c) filtering in the lag domain (i.e. by multiplying the variogram by some function) which is usually termed regularization in geostatistics. All of these methods produce very similar results and their differences are mainly of a methodological nature (Vanmarcke, 1983). Here we will adopt the lag domain approach (i.e. regularization) which is the one most frequently used in geostatistics. We adopt this approach because it is simple and intuitive and consistent with the variogram representation of variability used in the remainder of this paper.

The results presented here are based on standard regularization methods with a number of additional simplifying assumptions. In the case of support the variogram of an averaged process is calculated using the variogram of the point process and a filter function. The filter is represented by a square of side length  $a_{Supp}$ , which is the support. The square is the area over which the aggregation takes place. Different shapes of this area do not significantly affect the results of the regularization (Rodríguez-Iturbe and Mejía, 1974). In the case of spacing, the apparent variogram is approximated by the true variogram for lags larger than the spacing  $a_{Spac}$  and by a linear increase from the origin for shorter lags. This assumption is made because when estimating the empirical variogram there are only a small number of pairs of points for lags smaller than the spacing (Russo and Jury, 1987) and a straight line is the simplest approximation. Discussions of work related to this assumption are given in Russo and Jury (1987) and Gelhar (1993). In the case of extent, the apparent variogram is based on the true variogram for small lags (for  $\gamma(h) \leq \sigma_{app}^2$ ), and is constant and equal to  $\sigma_{app}^2$  for large lags. The apparent integral scale,  $I_{app}$ , is then calculated from the apparent variogram by using equation (3). Western and Blöschl (1999) showed that these assumptions are also reasonable for more general cases. More details on the method are given in Appendix A of Western and Blöschl (1999), Blöschl (1999), and in most geostatistical texts. Here we will show the results for two types of variograms: an exponential variogram (equation (2)), and a double exponential nested variogram of the form

$$\gamma = \sum_{i=1}^N \sigma_i^2 (1 - \exp(-h/\lambda_i)) \quad (8)$$

with  $N = 2$ ,  $\sigma_1^2 = 0.3$ ,  $\sigma_2^2 = 0.7$ ,  $\lambda_1 = 0.1$ , and  $\lambda_2 = 10$ . The variogram in equation (8) has two preferred scales, at 0.1 and at 10. From equation (3) we find that the integral scale of this variogram is  $I_{true} = 7.03$ . This type of variogram, with similar parameters, is sometimes used in hydrologic studies spanning a wide range of scales (e.g. Gelhar, 1993). It is important to reiterate that the filtering represents the change of scale (or scaling) and that the effect of this change of scale will be controlled by the ratio of the measurement scale (spacing, extent, support) and the process scale, and the ratio of the model scale (spacing, extent, support) and the process scale. It is therefore possible to use non-dimensional quantities such as  $a_{Spac}/I_{true}$ , i.e. the spacing normalized by the (true) process scale. Similarly, because of linearity, it is possible to non-dimensionalize the apparent variance,  $\sigma_{app}^2$  (which is the variance of the filtered process) by dividing it by the true variance,  $\sigma_{true}^2$ , and to non-dimensionalize the apparent integral scale (or correlation length),  $I_{app}$  (which is the integral scale of the filtered process) by dividing it by the true integral scale,  $I_{true}$ .

The results of the regularization analyses are shown in Figure 3. The top left panel of Figure 3 suggests that changing the spacing does not change the variance. For example, the SWE estimated from 10 and 1000 samples in a catchment will give the same variance, on average. This means that there is no bias introduced into the variance by a small number of samples but, of course, the estimate of the variance will be less reliable (large random error). No bias implies that the apparent variance is equal to the true variance and hence the ratio of the two quantities (plotted in the top left panel of Figure 3) is unity. Increasing the extent (top centre panel of Figure 3) increases the variance, and increasing the support (top right panel of Figure 3) decreases the variance. The effect of extent is consistent with the general observation that as we increase the size of the domain of interest, additional variability comes in. For example, when we start from the plot scale (say, 1 m<sup>2</sup>) the spatial variability of snow depth will be very small but as we increase the size of the domain (i.e. the extent) to a mountain range, the variability will increase drastically. For sufficiently large extents the apparent variance is equal to the true variance and hence their ratio approaches unity.

The effect of support (top right panel of Figure 3) is consistent with the general observation that aggregation always reduces variance for a random variable. For example, if we compare the SWE at a large number of plots in a region, the spatial variability will be quite large. However, if we aggregate the plot values to average values in a small number of catchments in the same region, the variability between the average catchment values will be much smaller. For sufficiently small supports the apparent variance is equal to the true variance and hence their ratio approaches unity. The reduction of variance with increasing



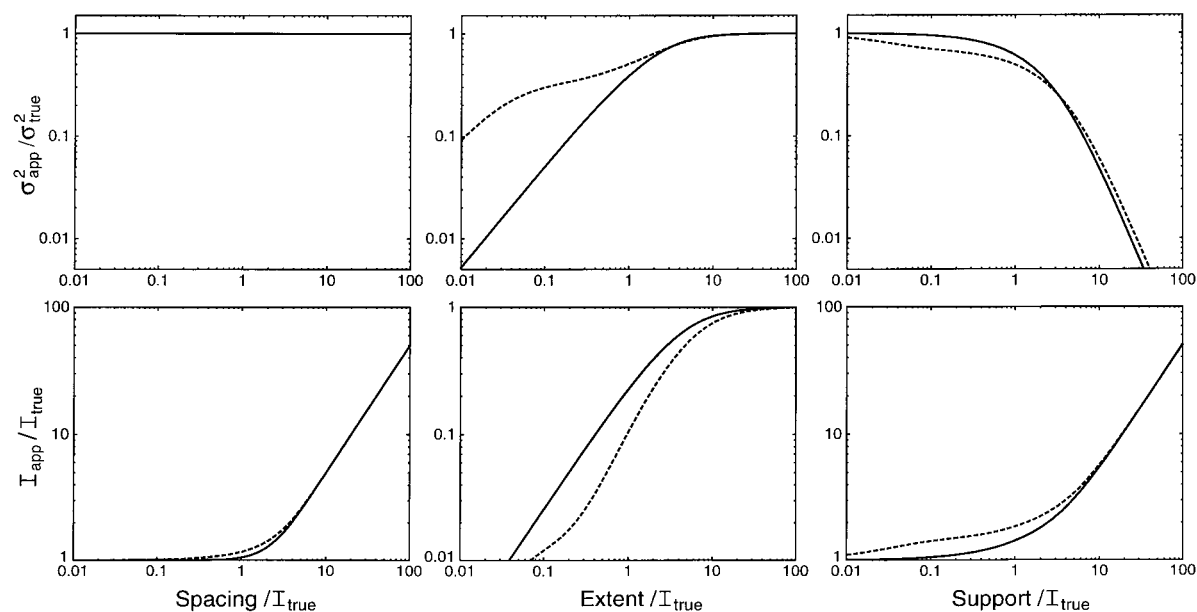


Figure 3. Effect of spacing, extent and support of a measurement or a model on biases in the variance and the correlation length.  $\sigma_{app}^2$  is the apparent variance;  $\sigma_{true}^2$  is the true variance;  $I_{app}$  is the apparent integral scale (or apparent correlation length);  $I_{true}$  is the true integral scale (or true correlation length). Based on geostatistical regularization methods (Western and Blöschl, 1999). Solid lines are for an exponential variogram (equation (2)) and dashed lines are for a double exponential nested variogram (equation (8)) with the parameters as in equation (8)

support has been recognized in many practical and more theoretical studies in various subdisciplines of hydrology including rainfall analysis (e.g. areal reduction factors) and regional flood frequency (see e.g. Blöschl and Sivapalan, 1997; Sivapalan and Blöschl, 1998).

Figure 3 also suggests that increasing the spacing, the extent, and the support will in general increase the apparent integral scale (lower panels). For example, a small number of samples of SWE in a large catchment analysed by geostatistical methods (equation (1)) may yield deceptively large correlation lengths, much larger than the correlation length of the underlying true snow patterns, because the spacing is large as compared to the process scale (i.e. true correlation length). On the other hand, correlation lengths estimated from snow lysimeter data at the plot scale are unlikely to capture the large scale variability of snow melt that is due to large scale topography and climatic conditions and, hence, will underestimate the correlation lengths of the underlying true pattern of snow melt in the region. This result is because the extent is small as compared to the process scale. Similarly, if we estimate correlation lengths from satellite data at a coarse resolution (i.e. large footprint or support), we will not be able to pick the small scale variability of snow cover patterns and hence we will overestimate the true correlation lengths.

It should be noted that Figure 3 shows the scale effect when changing only one of the scales of the scale triplet, assuming that the other two do not cause a bias (i.e. spacings are small, extents are large, and supports are small). The combined scale effect of two of the three components of the scale triplet has been examined in Blöschl (1999) and one example will be given later in this paper.

The effect of a change in scale on the mean,  $E(\cdot)$ , is not shown in Figure 3. This is because, given the assumption of a linearly aggregating process, a change of scale will not bias the mean. This result indicates that irrespective of the spacing, extent, or support of the data (or the model) the mean will always be unbiased, provided the process is linear. If the process model, or the measurement, does not aggregate linearly then the mean will be biased and may change with the scale. This is examined later in this paper. It

should also be noted that although the mean will not be biased for the case of linear averaging, the reliability (random error) of the mean will obviously change with the scale, but this is beyond the scope of this paper.

Let us consider two examples of how the effects depicted in Figure 3 can be used for practical applications. Consider, in the first example, a 10 km<sup>2</sup> catchment in which 100 point samples of snow depth have been taken. From these samples, a correlation length of about 200 m has been estimated by using equations (1) and (2). *Question:* (a) Is this estimate biased? (b) If so, what is the unbiased correlation length? (c) How many samples would be needed to obtain an unbiased estimate from the data? *Solution:* The solution is found by trial and error. Equation (4) gives  $a_{Spac} = 320$  m. If we assume that the true correlation length is 155 m,  $a_{Spac}/I_{true} = 2.06$  for which Figure 3 (bottom left panel) gives  $I_{app}/I_{true} = 1.29$ . This is a solution to the above problem as  $1.29 \cdot 155 \text{ m} = 200 \text{ m}$  which is equal to the apparent correlation length. This is a unique solution as, assuming, say,  $I_{true} = 180$  m gives an estimate of  $I_{app} = 220 \text{ m} > 200 \text{ m}$  while assuming  $I_{true} = 130$  m gives an estimate of  $I_{app} = 185 \text{ m} < 200 \text{ m}$ . The correlation length estimated from the samples is biased and the unbiased correlation length is about 155 m. For an unbiased estimate directly from the data (e.g. error smaller than 5%, i.e.  $I_{app}/I_{true} < 1.05$ ) Figure 3 (bottom left panel) indicates that  $a_{Spac}/I_{true} < 1$  which means that  $a_{Spac} < 155$  m and at least 420 point samples in space would be needed (equation (4)). It is interesting to examine the same example for the case of only 10 point samples for which  $a_{Spac} = 1000$  m and Figure 3 does not give a solution to the problem. While it is clear that the estimate will be substantially biased it is not possible to estimate the true correlation length. Clearly, there is not enough information in only ten samples. Similarly, there is no unique solution for apparent correlation lengths much smaller than the spacing which, again is due to the limited amount of information in the data.

Consider a second example of a remotely sensed image that gives some estimate of SWE. In a first step we are interested in the % snow covered area as estimated from this image. Assume that the mean SWE over the image is 360 mm, the true standard deviation (in space) is 300 mm, and the SWE is distributed according to a (truncated) normal distribution. Assume also that the true correlation length of SWE is  $I_{true} = 100$  m, and the pixel size (footprint) of the remotely sensed image is 250 m. *Question:* (a) What is the true snow covered area? (b) What % snow covered area would one predict from the remotely sensed image? *Solution:* (a) For the true SWE distribution, the mean is  $360/300 = 1.2$  times the standard deviation. For a standard normal distribution, the non-exceedance probability of  $P(Z < -1.2) = 0.12$ . This means that 12% of the land surface is snow free and 88% is snow covered. (b) From the remotely sensed image one would predict a variance that is smaller than the true variance. For  $a_{Supp}/I_{true} = 2.5$ , Figure 3 (top right panel) gives  $\sigma_{app,Supp}^2/\sigma_{true}^2 = 0.32$  which gives an apparent standard deviation of  $300 \cdot \sqrt{0.32} = 170$  mm. With this the mean is 2.1 times the standard deviation and for a standard normal distribution, the non-exceedance probability of  $P(Z < -2.1) = 0.02$ . This means that, from the remotely sensed image, one would predict that only 2% of the land surface is snow free and 98% is snow covered. Clearly, this misrepresentation will have a significant effect on estimates of the surface energy balance and evapotranspiration. As a side issue it is interesting to examine the apparent correlation length one would estimate from the images. For  $a_{Supp}/I_{true} = 2.5$  one gets  $I_{app}/I_{true} = 2.1$  from the bottom right panel of Figure 3, hence the apparent correlation length is 210 m rather than 100 m. It should also be noted that the numbers in this example depend on the assumption of a normal distribution.

### *Snow cover variograms*

So far we have implicitly assumed that (a) the true variograms of the snow cover variables are exponential (equation (2)) and (b) that the true correlation lengths exist and are known *a priori*. What is the data evidence on variograms of snow-related variables? Unfortunately, the variables of most interest (such as SWE and snowmelt) are usually collected at a few points only (e.g. Sommerfeld and Bales, 1993), hence, it is not possible to estimate variograms spanning many orders of magnitude in scale. Remotely sensed images are useful but some of the variables derived from them (such as SWE) may not always be reliable. Because of this, in this paper we examine snow covered area (SCA), i.e. a binary variable that is 1 for a snow covered pixel and 0 for a snow free pixel. A large number of snow covered area images are readily available and are

probably quite reliable. While we do not expect that the variograms of other snow-related variables will be identical with that of SCA, we do believe that SCA provides very useful insight into the nature of spatial variability (and variograms) of many snow-related variables, because the driving processes are closely interrelated. Patterns of SCA can in fact be interpreted as indicator images of SWE, representing different ‘thresholds’ of SWE at different times of the year. Indicator variograms and indicator geostatistics are widely used in other areas of hydrology and the geosciences in general where a continuous variable (such as SWE) is transformed into a binary variable by thresholding it (see e.g. Anderson, 1997; Western *et al.*, 1998). The correlation lengths derived from indicator variograms are closely related to the correlation length of the continuous variable and are of the same order of magnitude (Deutsch and Journel, 1997). However, of course it is not possible to estimate the variance of the continuous variable from the variance of the indicator variable. The variance of the indicator variable,  $\sigma_p^2$ , is always a function of the mean of the indicator variable. If  $p$  is the mean of, say, snow covered area (e.g.  $p = 0.3$ , i.e. 30% of the area is snow covered)

$$\sigma_p^2 = p \cdot (1 - p) \quad (9)$$

so the variance of the example is 0.21.

Sample variograms, estimated by equation (1), from four very different case studies are presented in Figure 4 as double logarithmic plots. The first set of variograms (Figure 4a) has been derived from a number

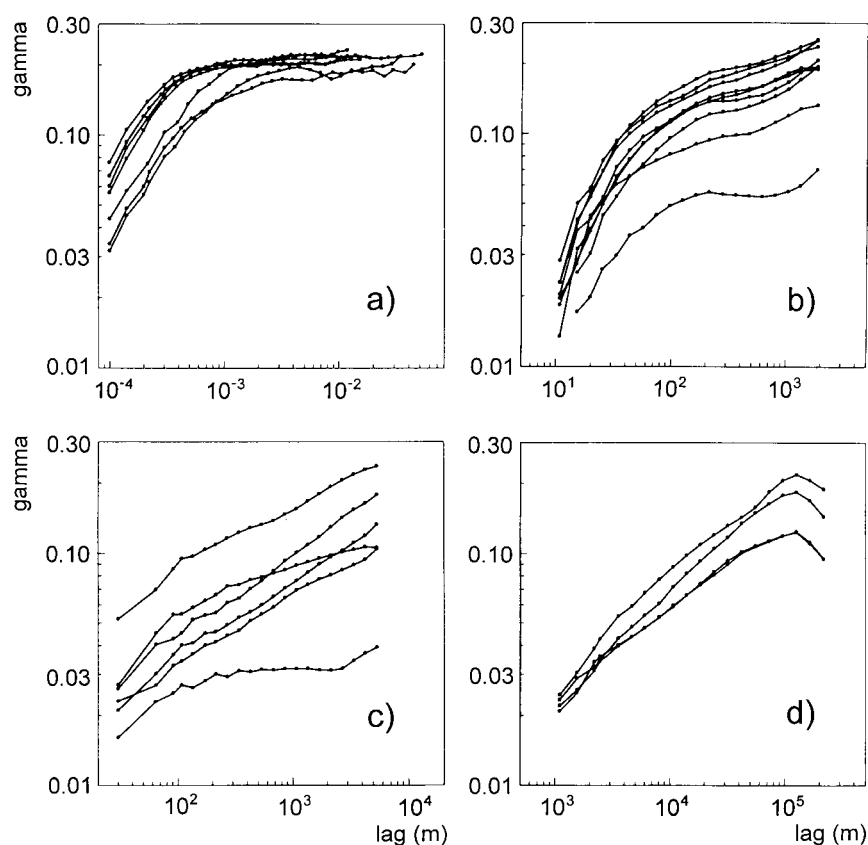


Figure 4. Variograms of snow covered area (SCA), i.e. a binary variable that is 1 for a snow covered pixel and 0 for a snow free pixel. (a) Thin sections of snow (eight images for different snow types, pixel size is 0.1 mm); (b) Kühtai aerial photographs (nine scenes in 1989, pixel size is 5 m); (c) Sierra Nevada TM images (six scenes in 1997, pixel size is 30 m); (d) Sierra Nevada AVHRR images (four scenes in 1998, pixel size is 1100 m)

of thin sections obtained in the laboratory by scanning images of snow crystals (R. Davis, *pers. comm.*). These are binary images where 1 is ice and 0 is void and the pixel size is 0.1 mm. Figure 4a shows that the variograms are all stationary, i.e. a correlation length exists and is on the order of 0.2 to 0.5 mm. The differences in the correlation lengths for the different images are mainly due to different types of snow. The shape of all variograms is close to exponential (equation (2)).

The second set of variograms (Figure 4b) has been derived from aerial photographs in the Kühltal catchment, Austria (Blöschl and Kirnbauer, 1992). The boundaries between snow covered and snow free areas were first digitized manually as vectors in the photographs and in a second step were gridded to a pixel size of 5 m. The variograms (Figure 4b) are probably not stationary but if we assume local stationarity (by neglecting the part of the variogram for lags larger than 500 m) we would get correlation lengths on the order of 100 m. The variance,  $\sigma_p^2$ , is a multiplicative factor in variograms (Figure 2a) and hence translates into the intercept in double logarithmic plots. The variograms in Figure 4b have similar slopes but different intercepts which means that the differences between the variograms are mainly due to differences in  $\sigma_p^2$  and hence due to differences in the % snow covered area (equation (9)).

The third set of variograms (Figure 4c) has been derived from Landsat Thematic Mapper (TM) images in the Sierra Nevada region (see Cline *et al.*, 1998a; Rosenthal and Dozier, 1996) with a pixel size of 30 m. The variograms are clearly not stationary as they do not flatten out at large lags over the range shown. Most of them can be closely approximated by a straight line in the double logarithmic plots which implies that they conform to a power law. The differences between the variograms are partly due to differences in the % snow covered area (equation (9)) which cause the vertical shifts in Figure 4c. However there are also differences in the slope which may be related to other factors such as changes in shading due to variable illumination angle in rugged terrain.

The fourth set of variograms (Figure 4d) has been derived from AVHRR images in the Sierra Nevada region (K. Elder, *pers. comm.*) with a pixel size of 1100 m. The variograms have a different shape than those in Figures 4a, 4b, and 4c. They are approximately stationary as they do not increase for lags larger than 100 km. However, they are not exponential and can be approximated by a straight line between lags of 3 km and 100 km. If we were to estimate correlation lengths from Figure 4d, they would be on the order of 30 km. The differences between the variograms in Figure 4d are mainly due to differences in the % snow covered area.

While the variograms in Figures 4a, 4b, 4c and 4d do not apply to the same date and the same location it is reasonable to assume that their general shape will be similar for other dates and locations. Their main difference then is the *scale* at which the snow cover data have been collected. Given the various biases introduced by the sensors in terms of their spacing, extent and support one wonders what the true variogram spanning more than nine orders of magnitude from the crystal to the regional scale would look like, i.e. the variogram that arises from a combination of the variograms at the four scales. There is an apparent inconsistency in the correlation lengths (Figure 4a:  $\lambda = 0.0005$  m; Figure 4b:  $\lambda = 100$  m; Figure 4d:  $\lambda = 30\,000$  m) and it would be interesting to find out whether it is possible to reconcile the variograms shown in Figure 4.

There are two philosophies in the literature on this issue. The first philosophy (e.g. Dagan, 1986; Neuman, 1990, 1993; Lovejoy and Schertzer, 1985) suggests that there is some universal continuous variogram that can be represented by a power law of the form

$$\gamma = a \cdot h^b \quad (10)$$

which is termed a variogram of a fractal and represents scale invariant (or self similar) behaviour. Equation (10) implies non-stationarity. This philosophy is consistent with data evidence when combining a large number of situations and case studies in hydrology. However, it is hard to understand what would be the actual processes giving rise to one continuous relationship. In other words, equation (10) relates the variability at the crystal scale to the variability at the regional scale and from a physical perspective one would not expect them to be related. The second philosophy (e.g. Gelhar, 1993, p. 295) suggests that there is a discontinuous variogram exhibiting steps of the form of equation (8). This variogram exhibits a number of preferred scales

(i.e. the  $\lambda_i$ ) each of which may represent one physical process. For example, the process scale of crystal growth may be  $\lambda_1 = 1$  mm, that of wind drift at hillslopes may be  $\lambda_2 = 1$  m, that of solar radiation effects at hillslopes of different aspects may be  $\lambda_3 = 100$  m, and that of different climatic conditions may be  $\lambda_4 = 10$  km. The combined nested variogram (equation (8)) then is a combination of the effects of the individual processes. It is locally stationary because it flattens out between the preferred scales. This interpretation is appealing because it is consistent with physical reasoning. For example, the thin sections in Figure 4a are stationary and this may be due to the presence of crystal growth processes only (such as equilibrium processes and kinetic processes) but no other (larger scale) processes (such as the formation of preferential flow fingers). On the other hand, as we increase the scale, larger scale processes are reflected in the snow cover data.

Perhaps one possible way of reconciling these two philosophies is by noting that the *ensemble mean* of a large number of nested variograms of the type of equation (8) with different sets of  $\lambda_i$  for different case studies may indeed combine to give an envelope that looks like the fractal in equation (10). For a particular case study it may be preferable to assume local stationarity at each of the scales mentioned as a working hypothesis. If the study does not span too many orders of magnitude, the assumption of a stationary variogram of the exponential type (equation (2)) or a similar variogram may therefore be appropriate which implies the existence of one correlation length. If the study spans more than, say, three orders of magnitude, a nested variogram (equation (8)) may be preferable as it is likely that a number of processes are operative at very different scales. For both types of variograms the effects of scale have been shown in Figure 3 (solid lines and dashed lines, respectively).

As a side issue it may be interesting to examine whether the slope of the variograms in the double logarithmic plots (i.e. parameter  $b$  in equation (10)) may be a useful parameter for characterizing the spatial snow cover variability. This is a potential alternative to the correlation length. It would also be interesting to know whether any relationships to the metamorphic state of the snow cover or other physical controls (such as the complexity of the terrain) can be detected. For the two dimensional case examined here, the parameter  $b$  is related to the Hurst exponent,  $H$ , by  $H = b/2$  and to the fractal dimension,  $D$ , by  $D = 3 - b/2$  which are parameters that are often used in studies of fractal variability (Klemeš, 1974; Feder, 1988; Klinkenberg and Goodchild, 1992). The exponent  $b$  is a measure of the ratio of large scale and small scale variability. If  $b$  is, say, 1.2 most of the variability occurs at large scales, and if  $b$  is, say, 0.1 most of the variability occurs at small scales. Straight lines have been fitted to the double logarithmic plots of Figure 4b (for lags larger than 50 m), Figure 4c, and Figure 4d (for lags between 3 and 100 km). The slopes of the fitted lines (i.e. the fractal exponents  $b$ , equation (10)) are shown in Figure 5 along with the proportion of snow covered area. The images are sorted according to the date they have been collected, i.e. from winter to summer as can be seen from the decreasing snow covered area which reflects the seasonal snow ablation. The exponent  $b$  does not vary much during the year. However, there is a slight tendency towards lower exponents at the end of the ablation period. This indicates a larger proportion of small scale variability as compared to large scale variability which may be a consequence of rocks and small scale topographic features having a stronger impact on the snow cover patterns towards the end of the ablation period. Similarly, image number 3 at Kühtai and image number 11 at the Sierra Nevada show slightly lower exponents. These images were collected shortly after snowfalls which again may cause more small scale features to be present in the patterns.

It is interesting that analyses of transects of snow depths reported in the literature tend to show piecewise fractal behaviour of the variogram for lags shorter than about 5 to 50 m, depending on the climate and the type of terrain (Shook and Gray, 1996; Sturm *et al.*, 1998). The integral scales in these studies were on the order of 2 to 20 m. Overall, the interpretation of the exponent  $b$  is not straightforward and it appears that more work needs to be done before it can be used as a useful parameter for characterizing spatial snow cover variability, particularly as far as its relation to the metamorphic state of the snow cover is concerned.

#### *Artefact or reality?*

As a final remark on the linear stochastic analysis we will draw attention to the subtleties of interpreting results of scale analyses. To illustrate the point we have resampled the Kühtai snow cover data used in

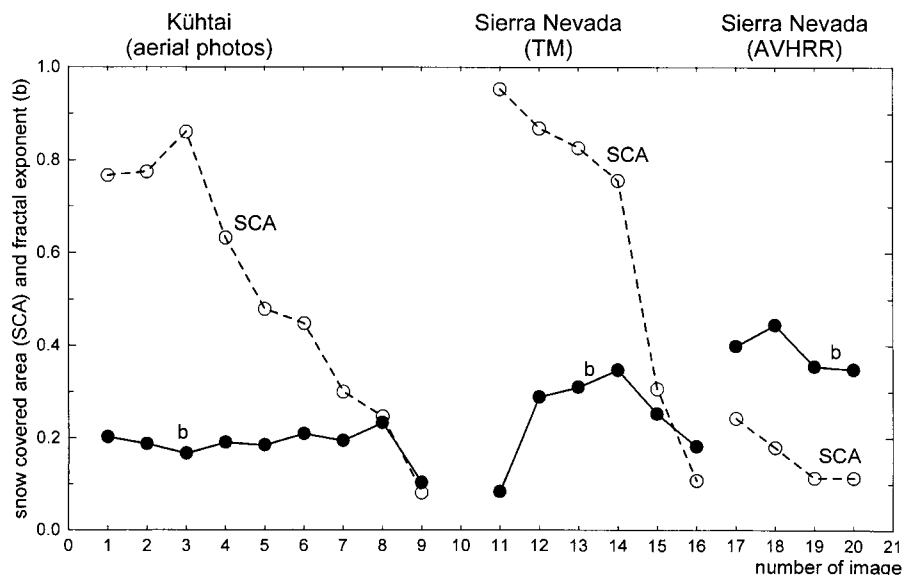


Figure 5. Fractal exponents  $b$  obtained by fitting a power law (equation (10)) to the variograms in Figure 4b (images 1–9), Figure 4c (images 11–16), and Figure 4d (images 17–20), and snow covered area (SCA)

Figure 4b (image number 8 in Figure 5). A square window of size  $a_{Ext}$  was placed at random in the catchment. From this window, 500 samples of snow cover were drawn randomly which were used to estimate the variogram and to fit an exponential variogram to it. The sizes of the windows were varied from 125 to 4000 m for the same centre of the window. As a final step, correlation lengths were plotted against window size. The results are shown in Figure 6a for four random locations of the window centres. Estimated correlation lengths tend to increase with window size. In other words, the apparent correlation scale increases with increasing measurement scale (extent). Similar results of scale dependent behaviour of correlation scales were reported by Gelhar (1993) for the case of hydraulic conductivities in aquifers. Specifically, the data of Figure 6.5 of Gelhar (1993) suggest that the correlation scales of conductivity tend to be on the order of 10% of the extent of the domain which is close to the results for the snow cover case shown here. Gelhar suggested that this behaviour is an indication of the nested nature of the true variogram which has the form of equation (8). However, this behaviour may also perfectly well be explained by the biases introduced by the sampling and the subsequent geostatistical analysis. In the resampling analysis we left the number of samples constant which implies a fixed ratio of the spacing and the extent. In Figure 6b we have expanded the analysis of Figure 3 by combining the cases of extent and spacing, i.e. we have calculated the apparent integral scale for an exponential variogram with varying extent and with the spacing fixed at a multiple of the extent. We used  $a_{Spac} = 0.1 \cdot a_{Ext}$  which implies 100 data points. Each of the three lines shown in Figure 6b relates to vastly different true correlation lengths (i.e. process scales,  $I_{true} = 0.01; 1.0; 100$ ). While the three lines are slightly different where the extent is close to the true correlation length  $I_{true}$ , from a global perspective all three lines centre around the dashed line that represents 10% of the extent. This means that the finding of correlation lengths to be 10% of the extent of the domain as suggested in Figure 6a and elsewhere in the literature may also be interpreted as an artefact of the method which does not allow inference of the true underlying hydrologic variability. In other words, if a wide range of scales is examined and only 100 data points or less are available, the apparent correlation length will always be on the order of 10% of the extent of the domain, irrespective of the shape and the correlation length of the true variogram. Obviously, there are serious inference problems (also see Gelhar, 1993, p. 330; Gallant *et al.*, 1994; and, for the temporal case, Klemeš, 1974) and one should, therefore, be wary of interpreting variability in the data as the true variability.

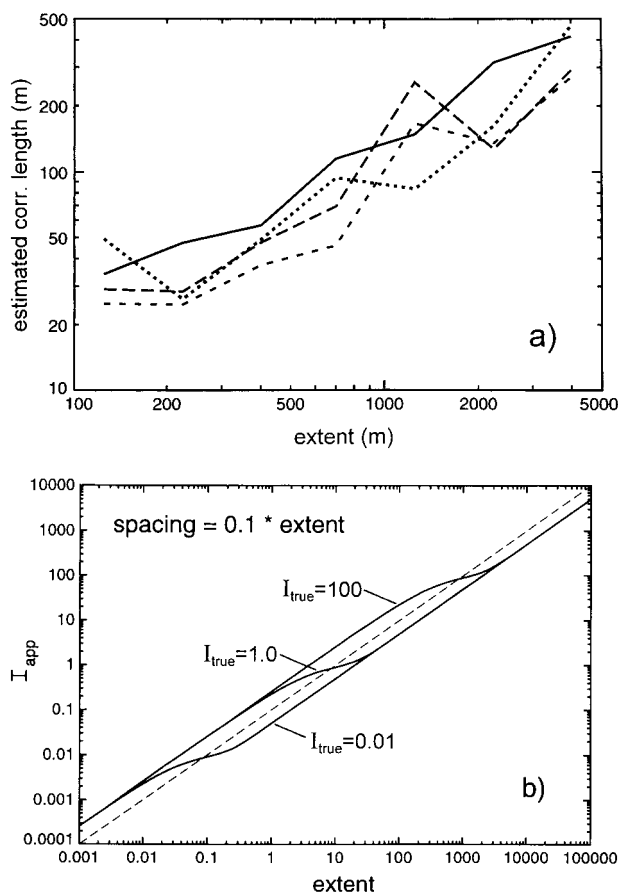


Figure 6. (a) Apparent correlation lengths estimated by resampling the Khtai snow cover data (Figure 4b and Figure 5, image number 8) plotted versus extent (i.e. size of a window). 500 data points in space have been used. The four lines correspond to four random locations of the window centre. (b) Joint effect of spacing and extent of a measurement on biases in the correlation length based on regularization methods using an exponential variogram (Western and Blschl, 1999).  $I_{app}$  is the apparent integral scale (or apparent correlation length);  $I_{true}$  is the true integral scale (or true correlation length). The spacing is 10% of the extent which is equivalent to 100 data points in space

## NON-LINEAR DETERMINISTIC ANALYSIS

### Framework

The non-linear analysis of spatial variability of snow processes clearly addresses a much wider class of cases than the linear analysis. Many processes in snow hydrology do not aggregate linearly and the associated variables are not conservative.

$$E(f(x)) \neq f(E(x)) \quad (11)$$

where  $E$  is the mathematical expectation,  $x$  is location and  $f$  is a function or a variable. The most important difference to the linear case is that a change in scale will have an effect on the mean,  $E(\cdot)$ , and hence different measurement or model scales can cause a bias in the mean. Strictly speaking, probably all processes associated with the formation, redistribution and depletion of the snow pack are non-linear as are the processes of meltwater movement in the pack and through the catchment. In many practical applications it is exactly the non-linearity that is of most interest. For example, if a number of snow courses are sampled with

the aim of identifying those locations that are more representative of the average over a catchment than others (e.g. Golding, 1974) it is clearly the non-linearity associated with the spatial organization of the snow cover that is examined. This is because if the SWE varied linearly, all locations in a catchment would be equally representative of the catchment mean. However, this is rarely the case (e.g. Yang and Woo, 1999).

A deterministic analysis of this non-linearity allows us to apply physical laws. Some of them (such as Darcy's law and laws associated with solar radiation) are relatively well known (e.g. Dozier, 1980) while others (such as laws associated with turbulent processes) are relatively poorly known or do not exist.

This sort of analysis is usually framed as a distributed physically based snow model (DSM) which allows incorporation of all the complexity of the physical processes that are deemed to be important and allows estimation of a range of variables that are of interest in snow hydrology, including SWE and melt water release from the snow pack (Leavesley, 1989; Kirnbauer *et al.*, 1994). To this end, the catchment is usually subdivided into numerical elements which are often square grid cells. The basic assumption in doing this is that the total spatial variability is split up into a small-scale part of variability *within* elements and a large-scale part of variability *between* elements (Smagorinsky, 1974; Kirnbauer *et al.*, 1994). The small scale variability within elements is also termed 'subgrid variability'. The large scale variability between elements is also termed 'element-to-element' variability. In a linear approximation, the total variability is the sum of the small scale variability within elements and the large scale variability between elements (see e.g. Isaaks and Srivastava, 1989)

$$\sigma_{total}^2 = \sigma_{within}^2 + \sigma_{between}^2 \quad (12)$$

In the framework adopted in the linear stochastic analysis of this paper (Blöschl, 1999),  $\sigma_{between}^2$  is simply  $\sigma_{app,Supp}^2$  as shown in Figure 3 (top right panel). DSMs deal with these two components in very different ways (see Figure 7; Kirnbauer *et al.*, 1994; and Blöschl and Sivapalan, 1995). The large scale processes (larger than the element size) are *explicitly* represented in the model, i.e. the variability is resolved. For the case of topography, explicit representation may be by different values of topographic elevation in different elements or by the sizes, shapes and slopes of the individual elements. The small scale processes (smaller than the element size) are *parameterized*, i.e. they are implicitly represented in the model by some lumped relationship. There are a number of reasons for using this approach. One of them is that, in most cases, the detailed small scale patterns of hydrologic processes within an element will not be known (or will even be unknowable; Beven, 1989, 1995) and hence a lumped representation as subgrid variability is appropriate and necessary.

To assess the importance of subgrid variability *vis à vis* total variability let us consider a DSM example and assume, as an approximation, that we can average the parameters linearly. Assume that the size of the model elements is 15 m and from detailed point samples the true correlation length of SWE is known to be 30 m. *Question:* (a) What is the variability of the average element SWE (i.e. the variability between elements) as

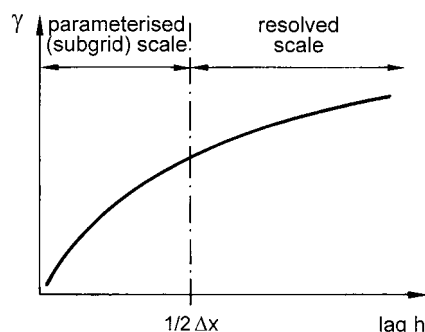


Figure 7. Hypothetical variogram to indicate that, by subdividing a catchment into model elements, distributed snow models split the total spatial variability into a small scale (subgrid) component which is parameterized and into a large scale component which is resolved explicitly.  $\Delta x$  is the element size



simulated by the model, assuming it is consistent with the true point scale variability? (b) What is the subgrid variability of SWE? (c) What is the correlation length of the patterns as simulated by the model? *Solution:* (a) For  $a_{Supp}/I_{true} = 0.5$ , Figure 3 (top right panel) gives  $\sigma_{app,Supp}^2 = 0.8\sigma_{true}^2$ . This means that the variability between elements is 80% of the total variability. (b) From equation (12) the subgrid variability is found as 20 % of the total variability. (c) Figure 3 (bottom right panel) suggests that  $I_{app,Supp} = 1.3 \cdot I_{true} = 40$  m. In other words, the change of scale considered in this example is not very important in terms of the variance and the correlation length. However, it should be noted that for many snow hydrologic processes non-linear effects caused by these differences including feedback effects may be vastly more important.

The representation of the spatial variability of snow-related processes between elements (i.e. the element-to-element variation) is usually done by some sort of spatial interpolation and there is a vast body of literature on this (e.g. Obled and Harder, 1979; Blöschl *et al.*, 1991; Elder, 1995; Davis *et al.*, 1995; including a number of papers in this journal issue). There is much less literature on subgrid variability and we will, therefore, touch on some methods of how to represent subgrid variability in DSMs.

### *Subgrid variability*

The main importance of representing subgrid variability accurately in DSMs probably stems from the feedback effects introduced by the non-linearity of the system. There are a number of approaches to quantifying the spatial variability of hydrologic processes within a grid cell (Blöschl and Sivapalan, 1995; Blöschl, 1996). These include:

- (a) the point value approach which assumes that the point scale equations at some point in the element suffice for an adequate description of the processes within the element. It is clear that this approach in essence neglects subgrid variability.
- (b) A second approach uses distribution functions rather than single point values. One example is the approach suggested by Luce *et al.* (1999) which extends a point mass and energy balance model by using a relationship between the basin average snow water equivalent and snow covered area to parameterize the subgrid variability. The relationship is similar to the 'snow cover depletion curves' in currently used empirical snowmelt models such as the National Weather Service River Forecasting System (Anderson, 1973).
- (c) A third approach uses effective parameters and assumes that the parameters and processes are uniform within each element and that the point equations apply to the whole element. This is a trivial problem for linear processes but very difficult for non-linear processes (e.g. Sivapalan and Wood, 1986; Blöschl and Sivapalan, 1995; Wen and Gómez-Hernández, 1996). In the non-linear case, there are two main questions; can the point scale equations be used to describe average element fluxes?, and if so, what is the scaling rule to obtain the effective parameters that are valid at the element scale (rather than at the point scale)? In principle, the effective parameters can be found by matching the flux obtained by the effective (average) parameter with the average flux in the element obtained with point scale parameters (flux matching). In practice, the effective parameters are often found by calibration.
- (d) A fourth possibility is a parameterization without explicitly resorting to the point equations. While I am not aware of any applications of this method in the context of spatial subgrid variability in DSMs, there is an excellent example in the time domain. Some point models of snow cover energy exchange explicitly simulate the snow surface temperature by modelling the vertical heat fluxes in the pack by a finite difference scheme (e.g. Blöschl and Kirnbauer, 1991). This type of model allows one to calculate the diurnal variations of the snow surface temperature that have important feedback effects on the surface energy balance. These models are usually run at a time step (temporal support) of one hour or less. Other models do not model the vertical heat fluxes in the pack and hence they are not able to calculate snow surface temperature, so it is usually set to 0 °C (e.g. Braun, 1985). These models are usually run at a time step (temporal support) of one day. For these models, the diurnal variations of the surface temperature

are subgrid variability (in the time domain). One way to parameterize this subgrid variability is by introducing a factor of refreezing. Energy losses from the snow pack are then multiplied by this factor. As this factor is usually on the order of 0.5, it reduces the night time heat losses of the pack. This produces a similar effect to the actual processes where the snow surface temperature drops during the night which gives a lower energy loss than would occur for a 0 °C surface temperature. The important point here is that there are no distribution functions and no effective parameters involved but an additional model component is introduced, i.e. a parameterization. It should also be noted that numeric models of atmospheric processes make wide use of parameterizations of subgrid variability (e.g. Houghton *et al.*, 1996).

#### *Optimum modelling scale*

There is a clear trade-off when selecting the size of the model grid cells which can also be seen from Figure 7 and equation (12). If we select a small grid size, we need to explain a lot of the variability explicitly (as element-to-element variability,  $\sigma_{between}^2$ ) which perhaps will be very difficult but we need to give little thought to subgrid variability. On the other hand, if we select a large grid size, representation of the element-to-element variability will be easier but, perhaps, representation of the small scale (subgrid) variability,  $\sigma_{within}^2$ , will be very difficult. It has been suggested that there may exist a grid size where this trade-off is at an optimum (Wood *et al.*, 1988). While in practice the selection of the grid size is often determined by practical considerations, such as data availability and the required resolution of the predictions, it is also interesting to ask whether there exists a scale which *generally* tends to be more appropriate as an element size than others.

One of the important questions to be addressed here is whether there are large scale processes that are clearly distinguishable from small scale processes. The desirable thing would then be to parameterize the small scale processes as subgrid variability and to explicitly represent (i.e. resolve) the large scale processes. There is no mix between the two processes which is a definite advantage for model building and calibration. Let us, again, consider an example from the time domain. There are snow processes at a seasonal time scale and these are clearly distinguishable from processes at the diurnal (within-day) time scale. It is wise to select the time step so as not to mix these two classes of processes. It would not be a good idea to use a 20 hour time scale. Clearly, the appropriate choice is either a time step small enough to resolve the diurnal variations (e.g. one hour) or to completely lump the diurnal variations into subgrid variability (e.g. time step of one day). Unfortunately, in the space domain, the choices are less obvious.

The body of literature (e.g. Wood *et al.*, 1988; Blöschl, 1996) addressing the question of an optimum model element size (i.e. support) generally starts from an analysis of spatial variability and explores whether there are clearly discernible scales of variability (i.e. small scale variability and variability at a much larger scale). An example of variability that shows two clearly discernible scales is that represented by the nested variogram in equation (8) which has preferred scales at  $\lambda_1 = 0.1$  and  $\lambda_2 = 10$  and little variability in between (i.e. the variogram flattens out between these two scales). In a second step these analyses generally suggest that the small scale variability may be attributable to one process, and the large scale variability may be attributable to another process and hence it may be prudent to choose the model grid size somewhere in between. In the example mentioned above it would be prudent to choose a model element size on the order of 1.

Spatial variability may be analyzed in a number of ways, for example by (a) a spectral analysis; (b) a variogram analysis, and (c) an examination of how the average values over an area change when increasing the size of that area. For linear processes these methods give essentially the same results (Vanmarcke, 1983), but for non-linear processes the first two may not be feasible and hence the third is generally used. However, as an approximation, the results of the latter can also be related to results of the first two methods. The idea of examining how the average values over an area change when increasing the size of that area was first conceived by Hubbert (1956) in the context of discussing the continuum assumption in groundwater flow theory, later used by Bear (1972) to define the Representative Elementary Volume (REV) as the order of magnitude where 'f (porosity) approaches smoothly a limiting value' (i.e. varies only smoothly with

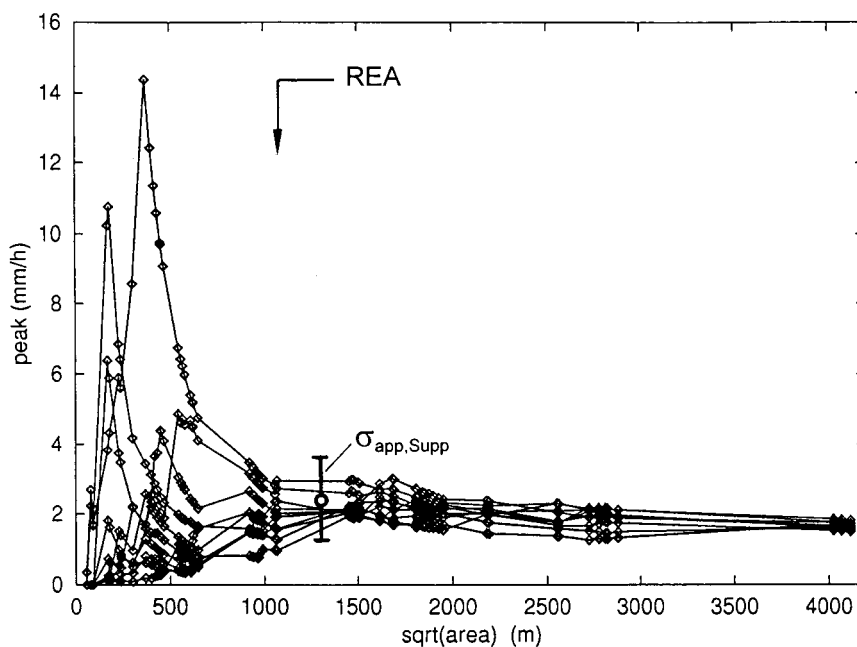


Figure 8. Peak flow simulated by a process-based rainfall-runoff model plotted versus subcatchment size for one set of nested catchments and ten realizations of spatially variable precipitation of one hour duration with correlation lengths of 125 m. From Blöschl *et al.* (1995). Compare Figure 3 top right panel

changing volume) and still later used by Wood *et al.* (1988) to define the Representative Elementary Area (REA) in catchment hydrology. The assumptions and results of Wood *et al.* (1988) have been re-examined a number of times (e.g. Famiglietti, 1992; Blöschl *et al.*, 1995; Fan and Bras, 1995). One example is given in Figure 8 where peak flows, as simulated by a distributed process-based runoff model (Grayson *et al.*, 1995), have been plotted versus subcatchment size for one set of nested catchments. The variability stems from assuming spatial random fields for the driving parameters (such as soil moisture and precipitation), from topographic variability, as well as from the non-linear overland and channel flow dynamics of the model. The different lines relate to different realizations of rainfall random fields.

From plots of the type shown in Figure 8 one would generally deduce an optimum model element size (i.e. REA) of about 1 km<sup>2</sup> as this is the size where peak flow smoothly approaches a limiting value. It is important to notice that, as an approximation, the results in Figure 8 can be related to results of the variance reduction in the linear case. Specifically, there is a close correspondence of the top right panel of Figure 3 (if plotted on a linear-linear scale) with Figure 8 where the area in Figure 8 corresponds to the support in Figure 3 and the variability between the realizations in Figure 8 corresponds to the apparent variance in Figure 3. Indeed, the decrease in variability in Figure 8 is mainly a consequence of the variance reduction due to increasing support which is controlled by the ratio of the support and the process scale. The most important driving parameter in Figure 8 is precipitation for which a correlation length (processes scale) of 125 m has been assumed in the simulations. Figure 3 indicates that the apparent variance gets small (10% of the true variance) at a support of about ten times the correlation length which closely resembles the results of Figure 8 where the variability smoothes out at about 1250 m. It is therefore likely that the results of analyses of the kind shown in Figure 8 are dominated by the assumptions made during the analysis and may, therefore, not be very useful for inferring the true behaviour of natural hydrologic variability. It should also be noted that the assumption of two clearly discernible scales (on which the concept of an optimum element size is based) is not born out in Figure 8 as there is no increase in variability at large scales (e.g. scales larger than 1000 m). This means that there is no large scale variability present in Figure 8. However, this large scale

variability is essential because it is exactly this large scale variability that is to be explicitly represented in distributed snow models by different values in different model elements.

Because of the reasons discussed above there is no clear evidence of the existence of an optimum element size in general. Indeed, Blöschl *et al.* (1995) have suggested that, an Arbitrary Elementary Area (AEA) of *any* size can be used. Clearly, this is the concept that is used in a number of disciplines related to snow hydrology such as hydrodynamic modelling and atmospheric modelling.

#### *Identifiability and index approach*

There are additional scaling problems with DSMs that are related to the complexity of the model (e.g. Obled, 1990; Grayson *et al.*, 1992, 1993). DSMs generally use a large number of complex equations each of which contains a number of parameters, and this complexity is multiplied by the large number of model elements which produces an excessively large number of degrees of freedom of the model. Unfortunately, it has been shown by various studies (e.g. Charbonneau *et al.*, 1981; World Meteorological Organisation, 1986; Loague, 1990) that predictions with *uncalibrated* catchment models tend not to be very accurate, no matter how much 'physics' these models contain. Because of this, there is always some element of calibration necessary for obtaining reliable DSMs. In some cases, this calibration is based on streamflow hydrographs, but alternative descriptions of the spatial snow cover processes of a catchment can give equally good fits of lumped catchment runoff (Blöschl *et al.*, 1994; Bathurst and Cooley, 1996). It may be very difficult to identify reliable model parameters of DSMs from runoff data alone.

Because of this, it has been suggested to use spatial snow cover data to calibrate and validate the models by comparing them to internal state variables of the models. Not only does this allow one to test the snow model directly but this also allows an assessment of its spatial predictive performance. Following the early paradigm of Blöschl *et al.* (1991), a number of studies in the recent literature have used snow cover patterns for evaluating DSMs (e.g. Davis *et al.*, 1995; Tarboton and Luce, 1996; Cline *et al.*, 1998a). While these studies generally concluded that the snow cover patterns are vastly superior to runoff in terms of assessing the DSMs, there is still a substantial degree of uncertainty associated with the predictions of DSMs which is mainly due to scaling problems.

In some instances, more parsimonious models are used for estimating the spatial patterns of snow-related variables in catchments to address the identifiability problem. These are generally index-type approaches that use the most important physical controls in a simple way (i.e. as indices), thereby maximizing the information that can be used, and at the same time minimizing the model complexity and hence problems associated with parameter identification. It is important to note that the indices are not purely statistical variables, rather they are derived from some (albeit simplified) process based reasoning (Moore *et al.*, 1991). Because of this, index approaches can be expected to perform better than purely statistical (black box) models, particularly in those situations that are different from the calibration period. One example in snow hydrology is the SWETREE model of Elder (1995). Figure 9 gives a sketch of the model structure in a typical configuration. The model is based on binary decision trees using regressions between SWE and indices for radiation and other controls. A pixel class is subdivided in steps, starting from the most important controls and proceeding to the less important controls. This approach was originally used for classifying remotely sensed images (Rosenthal and Dozier, 1996) and was later extended to ground based SWE measurements. However, as noted by Elder (1995), the advantage of a more parsimonious model structure comes at the cost of requiring a substantial data base for calibrating the model.

#### *Scale effects — comparison of non-linear and linear approaches*

The main purpose of DSMs is to simulate spatial patterns of snow-related variables. While, often, the spatial patterns are the main focus of interest, it is also interesting to see the scale effects on the spatial statistical *moments* of predictions from these non-linear models and how they are different from the scale effects in a linear stochastic analysis.

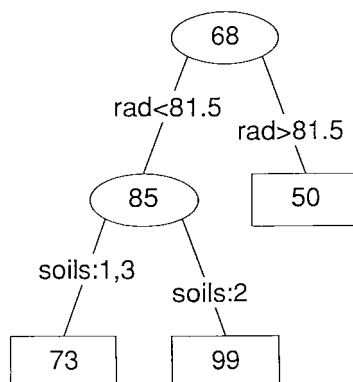


Figure 9. SWETREE regression tree for three snow water equivalent (SWE) classes for the 17–19 April, 1987 survey in the Emerald Lake watershed, California. Values in the ellipses represent the mean SWE value for all members of the subtree growing from that node. Values in the rectangular boxes represent the modelled SWE values from the field data satisfying the partition rules leading to that particular terminal node. Values are in centimetres of SWE. From Elder (1995)

In a first step (a) we therefore examine the effect of grid size (i.e. support of the model elements) on the spatial variances of snow water equivalent (SWE) between model elements simulated by a DSM. In a second step (b) we examine the effect of grid size on the mean SWE over a catchment.

(a) Effect on the variance  $\sigma_{app,Supp}^2$ : As an example we will present results from a sensitivity analysis of SWE as simulated by the DSM of Cline *et al.* (1998b) in the Emerald Lake watershed, California. Their model is based on energy balance calculations and remotely sensed data and simulates SWE on a regular grid. Cline *et al.* (1998b) started with a grid size of 30 m and degraded the model resolution to 500 m in steps. Figure 10a shows the (spatial) variance between model elements,  $\sigma_{app,Supp}^2$ , of SWE as calculated by the model for 21 April, 1993 plotted versus model grid size (i.e. support). There is a trend of decreasing variances with increasing grid size. It should be noted that for the largest grid size, only six pixels have been used to compute the variance, so the value may not be very reliable. Indeed, the decrease in variance is also consistent with a decrease in the range between the minimum and the maximum SWE in the catchment with increasing support. Table I of Cline *et al.* (1998b) gives, for the same date, ranges between the minimum and the maximum SWE in the catchment of 3.6, 2.27, 0.83, and 0.72 m for grid sizes of 30, 90, 250, and 500 m, respectively. The model used to calculate SWE is a non-linear model but it may be useful to compare the decrease in variance with the linear case. The corresponding linear case is shown in Figure 3 top right panel and also shows decreasing variances with increasing support, but an accurate comparison is difficult with the limited data in Figure 10a. Let us assume that the correlation length of SWE is  $I_{true} = 100$  m and the variance at the grid size of 30 m in Figure 10a is close to the true variance, so  $\sigma_{true}^2 = 0.292$  m<sup>2</sup>. With these assumptions, the largest grid size of 500 m gives  $a_{Supp}/I_{true} = 5$  and Figure 3 (top right panel) gives  $\sigma_{app,Supp}^2/\sigma_{true}^2 = 0.15$ . The variance of the largest grid size then is  $0.292 \times 0.15 = 0.044$  m<sup>2</sup> which is roughly what Figure 10a shows. However, the actual shape of the variance reduction is different from the linear case. This may be attributable to a number of reasons including the non-exponential shape of the variogram, the effects of non-linearity and, most importantly, the limitations of the small number of data points used for Figure 10a. However, it is clear that, overall, the scale effect of a reduction in variance when increasing the element size (i.e. support) is similar to the linear case.

(b) Effect on the mean  $E_{app,Supp}$ : As mentioned earlier in this paper, for linearly aggregating processes the statistical expectation,  $E_{app,Supp}$ , (or mean) will not change with the scale. However, for non-linearly aggregating processes there will be scale effects on the mean in the general case. This question is closely related to the question of effective parameters (see earlier in this paper) where the focus is on the bias in the mean when aggregating over one model element while now we examine the bias in the mean when aggregating over the entire catchment. There is a substantial body of literature in various subdisciplines of

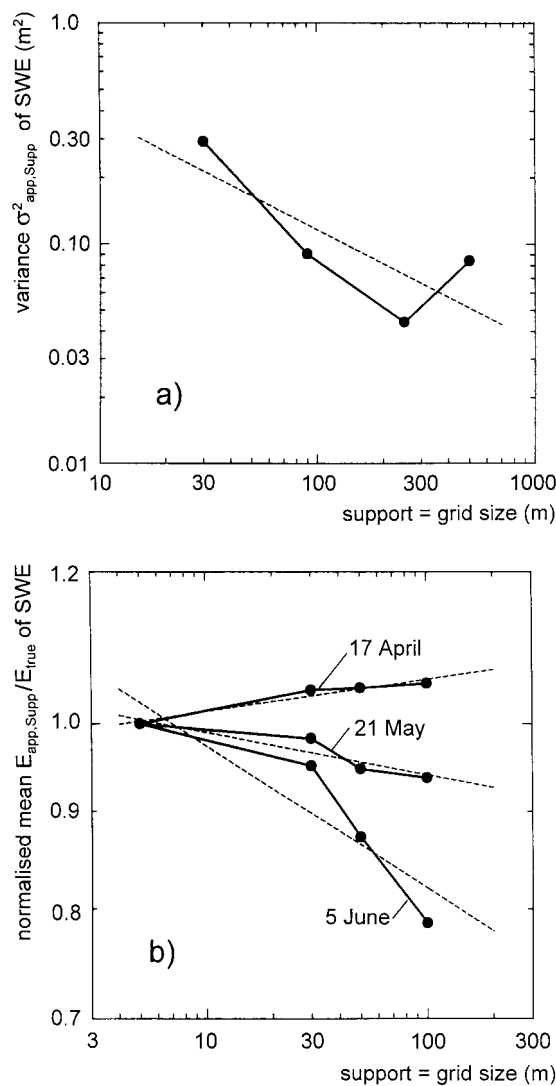


Figure 10. (a) Variance between model elements,  $\sigma_{app,Supp}^2$ , of snow water equivalent (SWE) as calculated by a distributed snow model for 21 April, 1993 in the Emerald Lake watershed, California, plotted versus model grid size (i.e. support). Compare with Figure 3 top right panel and Figure 8. The catchment area of the watershed is 120 ha. Plotted from information contained in Table I of Cline *et al.* (1998b). (b) Snow water equivalent (SWE),  $E_{app,Supp}$ , averaged over the Emerald Lake watershed calculated for various grid sizes (i.e. supports) by the SWETREE regression tree model. The average SWE,  $E_{app,Supp}$ , has been normalized by the average value for the 5 m grid resolution assuming this is close to the true value,  $E_{true}$ .  $E$  stands for the mathematical expectation. Plotted from information contained in Tables 23, 24, and 25 of Elder (1995)

hydrology on the effect of non-linearity on the aggregation of processes such as land surface-atmosphere interactions (e.g. Michaud and Shuttleworth, 1997; Becker *et al.*, 1999) and rainfall-runoff response (e.g. Goodrich *et al.*, 1997). Another important application with relevance to snow hydrology is the aggregation of digital elevation models (DEMs). For example, Band and Moore (1995) examined DEMs with different resolutions and found that the mean of the wetness index (where the wetness index is the logarithm of the ratio of specific contributing area and slope) increases with pixel size (i.e. support). Clearly, this is a consequence of the non-linearity present in the system. Slopes tend to decrease and specific

contributing areas tend to increase with pixel size. The net effect on the wetness index is a general increase in the mean with grid size.

For the snow cover case, results from the SWETREE regression tree model (Elder, 1995) are given here as an example. The SWETREE model is a non-linear model, hence one would not expect it to aggregate linearly and hence one would expect the mean SWE over a catchment to change with grid size (i.e. support). Figure 10b shows SWE averaged over the Emerald Lake watershed calculated for various grid sizes (i.e. supports) by the SWETREE model. In Figure 10b the average SWE,  $E_{app,Supp}$ , has been normalized by the average value for the 5 m grid resolution assuming this is close to the true value  $E_{true}$ .  $E$  stands for the mathematical expectation (equation (11)), i.e. the mean. Figure 10b indicates that at high snow accumulation levels (17 April, 1987) an increase in grid size (support) results in a minor increase in predicted SWE volume. Elder (1995) suggested that this minor effect is a consequence of linear snow free features being averaged into larger snow covered cells. This result implies that at peak accumulation the model averages close to linearly. Late in the melt season (21 May, 1987; 5 June, 1987), however, the grid size (support) becomes critical. Small snow patches are lost to averaging into large snow free cells and significant underestimation of the mean SWE results for large grid sizes. Elder (1995) suggested that the effect of scale is related to the size distribution of snow covered and snow free patches, as well as the geometry of the contiguous cells representing the snow covered area. This means that the main sources of non-linearity are related to the snow boundaries and the processes associated with the snow free/snow covered interface. This is consistent with the important role of snow covered area (SCA) in the representation of subgrid variability (Luce *et al.*, 1999). It is also important to note that SWE is a conservative quantity so, on physical grounds, it should average linearly. The scale effects seen in Figure 10b are a consequence of the non-linear averaging of the controls on SWE, such as terrain parameters and SCA, and how they are related to SWE in the SWETREE model.

Figure 10b indicates that the decrease of mean SWE with increasing support becomes more important towards the end of the season which implies that the non-linearity becomes more important when SWE is low. More generally speaking, this suggests that a high degree of non-linearity is associated with small values of the variable of interest. This finding is consistent with analyses of non-linearity in other subdisciplines of hydrology. For example, Merz and Plate (1997) found that the rainfall runoff processes in a small German catchment were close to linear for large events but were highly non-linear for small to medium sized events. Clearly, for large events rainfall exceeds the infiltration capacity significantly and hence the response is almost linear while for smaller events the rainfall is close to infiltration capacity and hence very non-linear. It is when rainfall intensity and infiltration capacity are similar that the non-linearity is at a maximum (Sivapalan and Wood, 1986). In snow hydrology, however, the spatial non-linearity is not very well understood and more work needs to be done to clearly identify the sources and the effects of non-linearity in spatial snow hydrologic modelling.

## CONCLUSIONS AND SOME FUTURE DIRECTIONS

The concept of scale can be used to quantify characteristic lengths of (a) a natural process (such as the correlation length of the spatial snow water equivalent (SWE) variability); (b) a measurement (such as the size of a snow density sample or the footprint of the satellite sensor), and (c) a model (such as the grid size of a distributed snow model). The different types of scales are denoted as process scale, measurement scale and model scale, respectively. Interpolations, extrapolations, aggregations, and disaggregations are viewed as a change in model scale and/or measurement scale.

In a first step we have examined, in a linear stochastic analysis, the effect of measurement scale and model scale on the data and the model predictions. It is shown that the ratio of the measurement scale and the process scale, and the ratio of the model scale and the process scale are the driving parameters for the scale effects. These scale effects generally cause biases in the variances and spatial correlation lengths of satellite images, field measurements, and simulation results of snow models. It is shown, by example, how these biases can be identified and corrected by regularization methods. At the core of these analyses is the variogram. For

the case of snow cover patterns, it is shown that it may be difficult to infer the true snow cover variability from the variograms, particularly when they span many orders of magnitude. Specifically, it may be difficult to infer preferred scales and stationarity but for practical applications, local stationarity may be a perfectly adequate assumption.

In a second step we have examined distributed snow models which are a non-linear deterministic approach to changing the scale. Unlike in the linear case, in these models a change of scale may also bias the mean over a catchment of snow-related variables such as SWE. There are a number of fundamental scaling issues with distributed models which include subgrid variability, the question of an optimum element size, and parameter identifiability. We give methods for estimating subgrid variability and point to potential avenues for parameterizing subgrid variability. We suggest that, in general, an optimum element size may not exist and that the model element scale may in practice be dictated by practical considerations such as data availability and the required resolution of the predictions. Model parameters may be more easily identified in spatial index approaches than in distributed physically based snow models but this is at the cost of requiring an extensive data set for proper calibration. The scale effects in distributed non-linear models can be related to the linear stochastic case. The biases in the mean caused by scale effects only occur in the non-linear case as they are a consequence of the non-linearity present in the natural system and/or the models. However, the reduction in variance with increasing element size (i.e. support) is similar to the linear case which gives some credence to generalizing the applicability of regularization methods, as an approximation, to non-linear cases.

It is likely that a number of areas of scale related research in snow hydrology will receive particular attention in the near future. The capabilities of obtaining spatial data sets of snow hydrologic quantities have been increased manifold in the past years through both satellite imagery and ground based networks (such as the US SNOTEL network; NRCS, 1998). There exist a number of outstanding data sets such as the snow depth transect data obtained by a sled mounted X-band FW-CW radar in the Kuparuk Basin in Alaska (Holmgren *et al.*, 1998) and in Antarctica (Sturm *et al.*, 1998). However, it is clear that this trend needs to continue. It is the patterns of variables related to snow processes that are likely to best advance the understanding of scaling effects in snow hydrology. Along with better information on the spatial patterns, there is a need for improved formal methods for assessing spatially distributed snow models based on spatial data. Work is progressing in this direction in various areas of hydrology (e.g. Western *et al.*, 1996). Such formal methods may be based on a range of statistical approaches including pattern recognition methods.

If we assume that random behaviour and linear aggregation of snow hydrologic variables apply, there are a range of methods available to assess scale effects and to perform interpolations, extrapolations, aggregations, and disaggregations in a consistent way. For the non-linear case this is vastly more difficult and hence most of the methods addressing the non-linear case are more *ad hoc*. It would be useful to extend the framework presented here to non-linear cases in the context of snow hydrology. Little is known about the feedback effects of subgrid variability introduced by the non-linearity of the system and yet they are probably extremely important in distributed snow models. Other subdisciplines of hydrology, such as subsurface hydrology, have an excellent track record in quantifying non-linear scale effects by stochastic methods in a consistent way (e.g. Dagan, 1986; Gelhar, 1993). Clearly, in the case of snow hydrology the emphasis of these methods needs to be on the spatial organization that is present in the natural snow cover environment.

In this paper we have examined the *biases* caused by the scale effects but of similar importance is the *reliability* of spatial estimates and how it is affected by a change of scale. With geographic information systems (GIS) and other spatial estimation methods readily available today, it is easy to interpolate between data points but it is much more difficult to assess how accurate these estimates are. It is, therefore, of great importance to assess the uncertainty of spatial estimates and to be explicit about uncertainty when presenting results of spatial interpolations, extrapolations, aggregations, and disaggregations. The assessment of spatial uncertainty is likely to be an area of increasing interest in snow hydrology in the near future.

While most of this paper has focused on physical snow processes, similar conclusions apply and similar methods are applicable to chemical and biological processes. Scaling problems may even be more acute for



chemical and biological processes, as in most applications chemical and biological data tend to be more limited than data about physical processes.

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