

# Statistical parametric maps in functional imaging: A general linear approach

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## Abstract

Statistical parametric maps are spatially extended statistical processes that are used to test hypotheses about regionally specific effects in neuroimaging data. The most established sorts of statistical parametric maps (e.g. Friston et al 1991, Worsley et al 1992) are based on linear models, for example ANCOVA, correlation coefficients and t tests. In the sense that these examples are all special cases of the *general linear model* it should be possible to implement them (and many others) within a unified framework. We present here a general approach that accommodates most forms of experimental layout and ensuing analysis (designed experiments with fixed effects for factors, covariates and interaction of factors). This approach brings together two well established bodies of theory (the general linear model and the theory of Gaussian Fields) to provide a complete and simple framework for the analysis of imaging data.

The importance of this framework is twofold: (i) Conceptual and mathematical simplicity, in that the small number of operational equations used are the same, irrespective of the complexity of the experiment or nature of the statistical model and (ii) the generality of the framework provides for great latitude in experimental design and analysis.

## Introduction

The aim of this paper is to describe and illustrate an implementation of the general linear model that facilitates a wide range of hypothesis testing with statistical parametric maps. This approach can be used to make statistical comparisons ranging from an unpaired t test to linear regression in the context of analysis of covariance. Its importance lies in (i) expediency; in that a single conceptual and algorithmic framework can replace the many currently employed; (ii) generality; in that experimental design is completely unconstrained (in the context of the general linear model) and (iii) portability; these linear models can be applied to any method of constructing a statistical parametric map.

Functional mapping studies (e.g. activation studies or comparing data from different groups) are usually analyzed with some form of statistical parametric mapping. Statistical parametric mapping refers to the construction of spatially extended statistical processes to test hypotheses about regionally specific effects. Statistical parametric maps (SPMs) are image processes with voxel values that have, under the null hypothesis, a known distributional approximation (usually Gaussian).

The power of statistical parametric mapping is largely due to the simplicity of the underlying idea: *Namely one proceeds by analyzing each voxel using any (univariate) statistical parametric test. The resulting statistics are assembled into an image, that is then interpreted as a spatially extended statistical process.* SPMs are interpreted by referring to the probabilistic behaviour of stationary Gaussian fields (e.g. Adler 1981). Stationary fields model not only the univariate (univariate means pertaining to one variable) probabilistic characteristics of the SPM but also any stationary spatial covariance structure (stationary means not a function of position). 'Unlikely' regional excursions of the SPM are interpreted as regionally specific effects (e.g. activations or differences due to pathophysiology). In activation studies regional or focal activation is attributed to the sensorimotor or cognitive process that has been manipulated

experimentally. This characterisation of physiological responses appeals to *functional specialization*, or segregation, as the underlying model of brain function. When comparing one group of subjects with another, a local excursion is taken as evidence of regional pathophysiology. When the physiological and cognitive (or sensorimotor) deficits are used to infer something about functional anatomy; this inference is usually based on a *lesion-deficit model* of brain organization. One could regard all applications of statistical parametric mapping as testing some variant of the functional segregation or lesion-deficit hypothesis.

The idea behind statistical parametric mapping is, of course, not new. Statistical parametric mapping represents the convergence of two earlier ideas; *change distribution analysis* and *significance probability mapping*. Change distribution analysis was a pioneering voxel-based assessment of neurophysiological changes developed by the St. Louis group for PET activation studies (e.g. Fox and Mintun 1989). This technique provided a mathematical underpinning for the powerful subtraction paradigm still employed today. Significance probability mapping was developed in the analysis of multichannel electrophysiological (EEG) data and involves the construction of interpolated pseudomaps of a statistical parameter. The fact that SPM has the same initials is not a coincidence, and represents a nod to its electrophysiological counterpart.

Unlike significance probability maps in electrophysiology, SPMs are well behaved in the sense they are approximated by stationary, spatially extended stochastic processes (due to the fact that neuroimaging samples uniformly and that the point spread function is stationary). This well behaved and stationary aspect of SPMs (under the null hypothesis) meant that theoretical advances were made, to a point where this area is growing rapidly and is an exciting part of applied spatial statistics. This development has been in the context of the theory of Gaussian fields (e.g. Friston et al 1991, Worsley et al 1992, Worsley et al 1993a, Worsley 1994, Friston et al 1994); in particular the theory of level-crossings (Friston et al 1991) and differential topology (Worsley et al 1992).

SPMs can be as diverse as the experimental design and univariate statistics used in their construction. Experimental designs in functional neuroimaging can be broadly divided into; (i) subtractive, (ii) parametric and (iii) factorial. Their application can be either to activation studies (time-series) or to group studies where each individual is studied once. Cognitive or sensorimotor subtraction is the best known example of the subtraction design in activation studies (e.g. colour - Lueck et al 1991, or higher cognitive function - Frith et al 1992). Categorical comparisons of different groups using SPMs represent another common application of statistical parametric mapping (e.g. depression - Bench et al 1992). Parametric designs include studies where some physiological, clinical, cognitive or sensorimotor parameter is correlated with physiology to produce an SPM of the significance of the correlation or regression. In activation studies this may be the time on target during a visuomotor tracking paradigm (Grafton et al 1993), performance on free recall (Grasby et al 1994), or frequency of stimulus presentation (Price et al 1993). In the analysis of different populations the parameter may reflect symptom severity (Friston et al 1992a) or some simple variable such as age (Martin et al 1992). Clearly the parameter being correlated can be continuous (e.g. time on target) or discrete (e.g. word presentation frequency). Factorial designs provide the opportunity to consider an interaction between the treatments or sorts of condition (two factors interact if the level of one factor affects the effect of the other. At its simplest an interaction is a difference in a difference). In activation studies this may be the interaction between motor activation (one factor) and time (the other factor), an interaction that provides information about adaptation (Friston et al 1992b). Another example of an interaction is between cognitive activation and the effects of a centrally acting drug (Friston et al 1992c, Grasby et al 1993). In summary there are many ways a hypothesis can be formulated and correspondingly there are as many sorts of SPMs.

All the examples in the previous paragraph used SPMs in conjunction with a test statistic (usually the t statistic or correlation coefficient) that represented a special case

of the general linear model. This begs the question 'is there a single general approach that could accommodate all the above examples (and more)?' The purpose of this work was to identify an instantiation of the general linear model that would do this.

This paper is divided into two sections. The first section deals with the theory behind the general linear model, the implementation and making statistical inferences. This section includes a brief review of how significant regional excursions (activation foci or other regionally specific effects) can be characterized and identified. The second section illustrates various applications of the approach, ranging from t tests to interaction terms in analyses of covariance. The applications are illustrated with an exemplary data set obtained during a word generation activation study of normal subjects.

## A general approach

### *The general linear model*

The general linear model for a *response* variable  $x_{ij}$  (such as rCBF) at voxel  $j = 1, \dots, J$  is:

$$x_{ij} = g_{i1}\beta_{1j} + g_{i2}\beta_{2j} + \dots + g_{iK}\beta_{Kj} + e_{ij} \quad 1$$

where  $i = 1, \dots, I$  indexes the observation (e.g. scan). The general linear model assumes the errors ( $e_{ij}$ ) are independent and identically distributed normally [ $N(0, \sigma_j^2)$ ]. For example in activation studies this means one is assuming an equal error variance ( $\sigma_j^2$ ) across conditions and subjects (but not from one voxel or brain structure to the next). Here the  $\beta_{kj}$  are  $K$  unknown parameters for each voxel  $j$ .

The coefficients  $g_{ik}$  are explanatory variables relating to the conditions under which the observation (e.g. scan)  $i$  was made. These coefficients can be of two sorts: (i) a *covariate* (e.g. global CBF, time, plasma prolactin level, ... etc) in which case eqn(1) is a familiar multivariate regression model or (ii) indicator-type or dummy variables, taking integer values to indicate the level of a factor (e.g condition, subject, drug ... etc) under which the *response* variable (rCBF) is measured. Mathematically speaking there is no distinction between these two sort of variables but we make the distinction for didactic reasons. Eqn(1) can be written in matrix form as a multivariate general linear model;

$$\mathbf{X} = \mathbf{G}\mathbf{B} + \mathbf{e} \quad 2$$

Here  $\mathbf{X}$  is a rCBF data matrix with elements  $x_{ij}$   $\mathbf{X}$  has on column for each voxel  $j$  and one row for each scan. The matrix  $\mathbf{G}$  is comprised of the coefficients  $g_{ik}$  and is called the design matrix. The design matrix has one row for every scan and one column for every effect (factor or covariate) in the model.  $\mathbf{B} = [\mathbf{B}_1 \mid \mathbf{B}_2 \mid \dots \mid \mathbf{B}_K]$  is the

parameter matrix where  $\beta_j$  is a column vector of parameters for voxel  $j$ .  $e$  is a matrix of normally distributed error terms. It may be noted that eqn(2) does not have a constant term. The constant term can be explicitly removed by mean correcting the data matrix or implicitly by adding a column of ones to  $G$ . For didactic purposes we assume the data  $X$  are mean corrected. Least squares estimates of  $\beta$  say  $b$ , satisfy the normal equations (Scheffe 1959 p9):

$$G^T G b = G^T X$$

if  $G$  is of full rank then  $G^T G$  is invertible and the least squares estimates are uniquely given by

$$b = (G^T G)^{-1} G^T X$$

$$\text{where } E\{b_j\} = \beta_j \text{ and } \text{Var}\{b_j\} = \sigma_j^2 (G^T G)^{-1} \quad 3$$

If the errors are normally distributed then the least squares estimates are also the maximum likelihood estimates and are themselves normally distributed (see Scheffe 1959).  $\text{Var}\{b_j\}$  is the variance-covariance matrix for the parameter estimates corresponding the  $j$ th voxel.

These simple equations can be used to implement a vast range of statistical analyses. The issue is therefore not so much the mathematics but the formulation of a design matrix ( $G$ ) appropriate to the study design and the inferences that are sought. This section describes one general approach that is suited to functional imaging studies.

The design matrix can contain both covariates and indicator variables reflecting the experimental design. Each column of  $G$  has an associated unknown parameter in the vectors  $\beta_j$ . Some of these parameters will be of interest [e.g. the effect of particular sensorimotor or cognitive condition or the regression coefficient of rCBF (the response variable) on reaction time (covariate)]. The remaining parameters will be of no interest



and pertain to confounding effects (e.g the effect of being a particular subject or the regression slope of voxel activity on global activity). Confounding here is used to denote an uninteresting effect that could confound the estimation of interesting effects (e.g. the confounding effect of global changes on regional activations). The two levels (*indicator vs covariate* and *interesting vs not interesting*) suggest that  $\mathbf{G}$  (and  $\mathbf{B}$ ) can be split twice into four partitions  $\mathbf{G} = [ \mathbf{G}_l | \mathbf{G}_c | \mathbf{H}_l | \mathbf{H}_c ]$  and similarly  $\mathbf{B} = [ \mathbf{B}_l^T | \mathbf{B}_c^T | \boldsymbol{\gamma}_l^T | \boldsymbol{\gamma}_c^T ]^T$  with estimators  $\mathbf{b} = [ \mathbf{b}_l^T | \mathbf{b}_c^T | \mathbf{g}_l^T | \mathbf{g}_c^T ]^T$ . Here effects of interest are denoted by the  $\mathbf{G}$  partitions and confounding effects of no interest by  $\mathbf{H}$  partitions. The subscripts l or c refers to the nature of the effect (level within a factor or a covariate). There is a fundamental distinction between effects that are interesting and those that are not, as will be seen below. The distinction between levels and covariates is not mathematically important but helps in understanding and describing the nature of the design matrix. Although there is no mathematical difference between treating the level and covariate separately and together, we actually deal with them as separate matrices in our software implementation. This helps to clarify things when reading the code and interacting with the programs. Each partition of the design matrix has one row for each scan and one column for each effect modelled by that partition. Using these partitions Eqn(2) can be expanded:

$$\mathbf{X} = \mathbf{G}_l \mathbf{B}_l + \mathbf{G}_c \mathbf{B}_c + \mathbf{H}_l \boldsymbol{\gamma}_l + \mathbf{H}_c \boldsymbol{\gamma}_c + \mathbf{e} \quad 4$$

where  $\mathbf{G}_l$  represents a matrix of 0s or 1s depending on the level or presence of some interesting condition or treatment effect (e.g the presence of particular cognitive component). The columns of  $\mathbf{G}_c$  contain the covariates of interest that might explain the observed variance in  $\mathbf{X}$  (e.g. dose of apomorphine or 'time on target').  $\mathbf{H}_l$  corresponds to a matrix of indicator variables denoting effects that are not of any interest (e.g. of being a particular subject or block effect). The columns of  $\mathbf{H}_c$  contain covariates of no interest or 'nuisance variables' such as global activity or confounding time effects  $\mathbf{B}_l$  are effects due to the treatments of interest (e.g. activation due to

verbal fluency).  $\beta_c$  are the regression coefficients of interest.  $\gamma_1$  are the effects of no interest (e.g. block or subject effects) and  $\gamma_c$  are the regression coefficients for the nuisance variables or confounding covariates.

To make this general formulation clear consider the model for an unpaired t test. In this instance  $\mathbf{G} = [\mathbf{G}_1]$ , where the elements of the column vector  $\mathbf{G}_1$  are 0 for all rCBF measurements in one group and 1 for the other group. A simple regression of reaction time on rCBF would be implemented by making  $\mathbf{G} = [\mathbf{G}_c]$  where  $\mathbf{G}_c$  is a column vector containing the reaction time data. The randomized block design ANCOVA implemented by the MRC SPM software corresponds to  $\mathbf{G} = [\mathbf{G}_1 | \mathbf{H}_1 | \mathbf{H}_c]$  where  $\mathbf{G}_1$  specifies the activation condition, the  $\mathbf{H}_1$  account for subject (block) effects and  $\mathbf{H}_c$  is a column vector of confounding global CBF covariates. The point to be made here is that nearly every conventional statistical design is a special case of eqn(4) or in other words *most linear parametric analyses can be implemented with eqn(4)*.

In this paper scaling the images to remove global or whole brain effects is considered a pre-processing step. If we tried to incorporate a geometric scaling into the statistical model we would end up with a model that was more complicated than those considered here.

#### *Experimental design - the forms for $\mathbf{G}_1$ and $\mathbf{H}_1$*

In functional imaging one measures some physiological variable, usually hemodynamic (e.g. rCBF) in a number of conditions (levels of a factor). In many instances the same conditions are measured in a series of blocks; for example in PET, baseline and activations (conditions or factor levels) are repeated in a number of subjects (blocks or plots). In fMRI different conditions are usually repeated serially in blocks over time.

The situation gets more complicated if one repeats the activation study under different conditions or in different subjects. The designs appropriate for this situation are called split-plot designs [a good reference for psychology applications is Winer 1971) and see Worsley et al (1991) for an example in the PET literature using region of interest

analysis]. There are some complications that can arise when using split-plot designs, however the situation is simplified if the block (or subject) effects are always included thereby restricting the analysis to within-block effects. This is the approach taken here.

An example of this sort of experimental design, in PET, would be two verbal fluency studies, one in a series of normal subjects and one in a group of schizophrenic patients, or a psychopharmacological activation study in two groups of normal subjects studied with and without an appropriate antagonist. An example in fMRI would be the repetition of a blocked series of sensorimotor conditions in a different subject. In these examples the design is not completely blocked. In other words all conditions within and between studies are not repeated in the same block (e.g. subject). For example one cannot perform a verbal fluency activation study in a subject, give the subject schizophrenia, and then repeat the activation study. This partial blocking problem suggests the following form for  $\mathbf{G}_1$  and  $\mathbf{H}_1$ : Let the experiment be divided into a series of *studies*. In each of the  $s = 1, \dots, S$  studies  $n_s$  conditions are measured in all of  $m_s$  subjects (or more generally blocks). There is however no requirement for the number of conditions or subjects (blocks) within each study to be the same. This design is, we consider, the most accommodating for functional imaging experiments. One can think of this design as allocating  $n_s$  conditions to  $m_s$  blocks in any permutation allowed. The column position of the '1' in the  $i$ th row of  $\mathbf{G}_1$  denotes which of the  $n_s$  conditions the  $i$ th scan was obtained under and the position of the column with a '1' in the  $i$ th row of  $\mathbf{H}_1$  denotes the subject (block) in which that measurement was made.

The suggested terminology, adopted in this paper, divides the *experiment* into *studies*. Each study comprises one or more scans on one or more *subjects* (more generally blocks). Each subject is scanned in one or more *conditions*. *Conditions* can be variously classified and indeed may include several *treatments* [e.g. entreating subjects to perform a series of psychological tasks (cognitive treatments) with and without a drug (pharmacological treatments)]. A single level of any cognitive or sensorimotor treatment is a *task* and may be repeated in a number of conditions; but note that the first time a task is administered is one condition and the second time is

another condition. The design matrix depends only on *studies*, *subjects* and *conditions*. The effects of a specific *task* are assessed by comparing one set of conditions with another using contrasts, as discussed in later sections.

This one-way layout does not explicitly model interaction terms or effects due to study. This simplicity is deliberate and allows the investigator to test for the [simple] main effects of condition and interactions between condition effects and study *post hoc* (by testing for the appropriate effects on the  $n_s$  condition specific parameters  $\mathbf{B}_1$ ). The alternative would be to model these effects explicitly in  $\mathbf{G}$ . The problem with this approach is that the effect of interest would only be assessed in an omnibus sense. For example imagine a pharmacological activation (with  $P$  scans or conditions) performed in three groups, a normal, a schizophrenic and a depressed group. The interaction between condition and study (i.e. a regionally specific differential response to the drug challenge, across the three diagnostic groups) could be assessed using the ratio of variances (see below) but there would be no information about which group differed from which, or about the nature of the effect (augmented or blunted response). All these pairwise, direction-specific comparisons could be made *post hoc* if the experiment was treated as  $3P (n_s)$  distinct conditions. The proposed layout does exactly this.

A note of caution here: The multiple study (or split-plot) approach is proposed primarily for the analysis of interactions between condition and studies (and main effects of condition). Testing for the main effects of study is not so straightforward. This is because the estimates of the error terms are based on a within-block analysis and may not be appropriate for comparing between-block effects such as those due to study (See Winer 1971).

The solution implicit in eqn(3) requires that  $\mathbf{G}$  be of full rank or non-singular. This is guaranteed for the covariates, assuming all the covariates are linearly independent; however it is sometimes the case that the  $\mathbf{G}_1$  and  $\mathbf{H}_1$  are not linearly independent. This is dealt with by applying constraints to the design matrix. In the current context this is effected by constraining all the block effects (effects not of interest) to sum to zero.

This is equivalent to treating the uninteresting effects as residuals about the effects of interest.

*Adjusting for the confounding effects of no interest*

It is often wise to report physiological changes from a particular voxel in order to indicate the size of the physiological effect. This is best demonstrated by reporting the data (e.g. rCBF) after adjusting for the confounding effects of nuisance variables (e.g. global activity) and other spurious effects (e.g. block effects). After estimating  $\mathbf{B}$  the adjusted data  $\mathbf{X}^*$  are given by discounting the effects of no interest (i.e. in mean corrected form):

$$\mathbf{X}^* = \mathbf{X} - [\mathbf{H}_1 | \mathbf{H}_c] \cdot [\mathbf{g}_1^T | \mathbf{g}_c^T]^T$$

5

$[\mathbf{H}_1 | \mathbf{H}_c]$  represents the uninteresting or confounding parts of the design matrix (obtained by stacking the uninteresting factor level partition and the uninteresting covariates partitions side by side).  $[\mathbf{g}_1^T | \mathbf{g}_c^T]^T$  represents the corresponding parameter estimates. The  $\mathbf{g}_1$  and  $\mathbf{g}_c$  are linear estimators of  $\gamma_1$  and  $\gamma_c$ , the uninteresting components of  $\mathbf{B}$ . The adjusted *mean* condition affects are given by the elements of  $\mathbf{b}_1$  and the regression coefficients by  $\mathbf{b}_c$ .

*Statistical inference - Omnibus or overall effects at each voxel*

In this section we address statistical inferences about the effects of interest (condition and covariates of interest). We also clarify the relationship between statistical parametric mapping (i.e. massive univariate testing) and multivariate analysis.

The omnibus significance is assessed by testing the null hypothesis that including the effects of interest does not significantly reduce the error variance. This is equivalent to testing the null hypothesis that  $\mathbf{B}_1$  and  $\mathbf{B}_c$  are zero. In the sense that this inference does not address any specific condition (or covariate) it is referred to as an omnibus test.

Note that in contradistinction to the use of 'omnibus' in reference to tests that pertain to the whole image (e.g. the gamma-2 of Fox et al 1988) 'omnibus' is used here in a univariate sense for a particular voxel [this use of omnibus is preferred because detecting a significant regional effect (see below) is also an implicit confirmation that an effect exists somewhere in the image; i.e. in the old omnibus sense].

The null hypothesis that the effects embodied in  $\mathbf{G}_1$  and  $\mathbf{G}_c$  are not significant can be tested in the following way. The sum of squares and products due to error  $\mathbf{R}(\ )$  is obtained from the difference between the actual and estimated values of  $\mathbf{X}$  (  $\ )$  is the alternative hypothesis to the null hypothesis and includes the effects of interest):

$$\mathbf{R}(\ ) = (\mathbf{X} - \mathbf{G} \cdot \mathbf{b})^T (\mathbf{X} - \mathbf{G} \cdot \mathbf{b}) \quad 6$$

The error sum of squares and products under the null hypothesis  $\mathbf{R}(\ 0)$  i.e. after discounting the effects of interest ( $\mathbf{G}_1$  and  $\mathbf{G}_c$ ) are given by:

$$\mathbf{R}(\ 0) = (\mathbf{X} - [\mathbf{H}_1 \mid \mathbf{H}_c] \cdot [\mathbf{g}_1^T \mid \mathbf{g}_c^T]^T)^T \cdot (\mathbf{X} - [\mathbf{H}_1 \mid \mathbf{H}_c] \cdot [\mathbf{g}_1^T \mid \mathbf{g}_c^T]^T) \quad 7$$

Clearly if  $\mathbf{H}_1$  and  $\mathbf{H}_c$  do not exist this simply reduces to the sum of squares and products of the response variable ( $\mathbf{X}^T \mathbf{X}$ ). At this point we could (in principle) proceed in one of two directions. The first direction would be to test the omnibus significance of all effects of interest *over all voxels*. This would correspond to a multivariate analysis of variance or covariance (MANOVA or MANCOVA). Although many standard texts deal with MANOVA and MANCOVA separately there is no distinction in the context of the current analysis. The second approach would be to test for omnibus significance over effects of interest in a univariate sense at each voxel (c.f. ANOVA or ANCOVA at each voxel). The second is employed by statistical parametric mapping and preserves the regional specificity of the omnibus test at the cost of not being able to

extend the 'omnibusness' to the whole brain (i.e. a multivariate omnibus test). The multivariate omnibus significance would be tested with a single statistic, for example:

$$= \frac{|\mathbf{R}(\ )|}{|\mathbf{R}(\ 0)|} \quad 8$$

where  $\lambda$  is Wilk's statistic (known as Wilk's Lambda). A special case of this test is the Hotelling  $T^2$  test (see Chatfield and Collins 1980). However there is a problem here; namely  $\mathbf{R}(\ )$  and  $\mathbf{R}(\ 0)$  are both singular (meaning  $\lambda$  is undefined). This is a simple result of having more voxels than scans. This is one reason statistical parametric mapping adopts a mass univariate approach (mass univariate analyses implemented in parallel at each voxel). For a single voxel  $j$  eqn(8), after appropriate transformations reduces to the univariate ratio of variances (see Chatfield and Collins 1980 for details):

$$F_j = \frac{(r/[r_0 - r]) \cdot [R_j(\ 0) - R_j(\ )]}{R_j(\ )} \quad 9$$

where  $R_j(\ )$  corresponds to the  $j$ th element on the leading diagonal of  $\mathbf{R}(\ )$  and similarly for  $R_j(\ 0)$ .  $F_j$  is distributed according to the F distribution with degrees of freedom  $r_0 - r$  and  $r$ . Generally  $r = I - \text{rank}(\mathbf{G})$  and  $r_0 = I - \text{rank}(\begin{bmatrix} \mathbf{H}_1 & \mathbf{H}_c \end{bmatrix})$  where  $I$  is the total number of scans. If the data have been mean corrected and  $\mathbf{H}_1$  has not, it is necessary to remove an extra degree of freedom from  $r_0$ . The  $F_j$  can be displayed as an image to create an SPM{F} directly testing the overall significance of all effects designated 'of interest'. In practice SPM{F}s are seldom employed as direct tests of hypotheses but this form of omnibus testing is very useful for selecting subsets of voxels that are used in some further analysis [e.g. singular value decomposition or principal component analysis. See Friston et al (1993)].

#### *Statistical inference - specific effects at each voxel*

In the previous section the F ratio of variance was used to make some inference about the effects of *all* conditions and covariates of interest. In this section the significance of

*specific* effects is examined. This is effected with the t statistic using linear compounds or contrasts of the parameter estimates  $\mathbf{b}_j = [b_{1j}, b_{2j}, \dots]^T$ . For example if we wanted to test for activation effects between conditions one and two then we would use the contrast  $\mathbf{c} = [-1 \ 1 \ 0 \ 0 \ \dots]$ . The significance of a particular linear compound of effects at voxel  $j$  is tested with:

$$t_j = \frac{\mathbf{c} \cdot \mathbf{b}_j}{s_j}$$

where the standard error for voxel  $j = s_j$  is estimated by  $s_j^2$  [c.f. eqn(3)]:

$$s_j^2 = \frac{R_j(\mathbf{c})}{r} \mathbf{c} \cdot (\mathbf{G}^T \mathbf{G})^{-1} \cdot \mathbf{c}^T \quad 10$$

$t_j$  has the Student's t distribution with degrees of freedom  $r$ . When displayed as an image the  $t_j$  constitute an SPM $\{t\}$  and represent a spatially extended statistical process that directly reflects the significance of the profile of effects 'prescribed' by the contrast  $\mathbf{c}$ .

### *Statistical inference - specific effects - over the entire SPM*

In this section we address the problem of how to interpret the SPM $\{t\}$  in terms of probability levels or p-values. The problem here is that an extremely large number of non-independent univariate comparisons have been performed and the probability that any region of the SPM will exceed an uncorrected threshold by chance is relatively high. In recent years advances have been made that have solved this problem. These advances have focussed on choosing thresholds that render the chance probability of finding an activation focus, over the entire SPM, suitably small (e.g. 0.05). The threshold can be of two types (i) a critical height that the region has to reach or (ii) a critical size (above some threshold) that a region must exceed before it is considered significant. Another way of looking at these approaches is to consider that a local excursion of the SPM (a connected or contiguous subset of voxels above some



threshold) can be characterized by its maximal value ( $Z$ ) or by its size ( $n$  - the number of voxels that constitute the region). Both these simple characterizations have an associated probability of occurring by chance over the volume of the SPM. These two probabilities form the basis for making a statistical inference about any observed regional effect:

The analyses behind the distributional approximations for  $Z$  and  $n$  derive from the theory of continuous, strictly stationary, stochastic Gaussian random fields (Gaussian here refers to the multivariate Gaussian probability distribution of any subset of points, *not* to the autocorrelation function). To simplify the analysis the  $SPM\{t\}$  is transformed to a  $SPM\{Z\}$  using a probability integral transform or other standard device. Strictly speaking this univariate transformation does not make a  $t$ -field into a Gaussian field unless the degrees of freedom of the  $SPM\{t\}$  are very high. In what follows we assume that we are dealing with reasonably high degrees of freedom and that the  $SPM\{Z\}$  is a reasonable lattice representation of an underlying continuous Gaussian field. Analytical expressions for the distributional approximations of  $t_{\max}$  (the largest  $t$  value) have now been established (Worsley et al 1993, Worsley 1994) however approximations for  $n_{\max}$  (the size of the largest region) have not.

*The p-value based on Z, the largest value in the region*

The probability of getting at least one voxel with a  $Z$  value of say height  $u$  or more, in a given  $SPM\{Z\}$  of volume  $S$  [ $P_S(Z > u)$ ] is the same as the probability that the largest  $Z$  value is greater than  $u$  [ $P(Z_{\max} > u)$ ]. This is the same as the probability of finding at least one region above  $u$ . The central tenet here relies on the fact that the probability of getting at least one region above  $u$  and the expected number of regions tend to equality (at high values of  $u$ ).

$$P_S(Z > u) = P(Z_{\max} > u) = P(m > 1) = E\{m\}$$

Where  $m$  is the number of regions. The problem therefore reduces to finding the expected number of foci at  $u$ . An analysis was presented in Friston et al (1991) that showed how this expectation could be identified using the theory of Gaussian fields. This analysis approximated activation foci with suprathreshold ellipsoid regions. Subsequently the Euler characteristic (the number of blobs minus the number of holes) has been proposed as an approximation to the number of foci. The Euler characteristic was introduced in a key paper by Worsley et al (1992) that also established a formal link between the theory presented in Friston et al (1991) and earlier work on the expected number of maxima (see Adler 1981) [the expressions derived on the basis of ellipsoid regions (Friston et al 1991) and those based on maxima (Adler 1980) were shown to have, asymptotically, the same form but differ by a factor of  $\pi/4$  at infinitely high thresholds (Worsley et al 1992)]. The important issue here is that whether one uses the number of foci, the Euler characteristic or the number of maxima, very consistent results are obtained. At high thresholds the number of regions, the number of maxima and the Euler characteristic all tend to the same value. The Euler characteristic is more amenable to mathematical analysis than the earlier formulations and has led to extensions to SPM{t} and SPM{F} (Worsley et al 1993a, Worsley 1994). For simplicity we will work with the number of maxima (Hasofer 1978):

$$P(Z_{\max} > u) \approx E\{m\} \approx \frac{S}{W^D} (2\pi)^{-(D+1)/2} W^{-D} u^{D-1} e^{-u^2/2} \quad 12$$

$W$  is a measure of smoothness and is related to the full width at half maximum (FWHM) of the SPM's "resolution". Equivalently  $W$  is inversely related to the number of "resolution elements" or Resels ( $R$ ) that fit into the total volume ( $S$ ) of the  $D$ -dimensional SPM. ( $R = S/W^{D+1}$ ).

In practice  $W$  can be determined directly from the effective FWHM if it is known when  $W = \text{FWHM} / (4 \log_e 2)$  or estimated post hoc using the measured variance of the SPM{Z}'s first partial derivatives:

$$W = \prod_{i=1}^D \text{Var}\{ \text{SPM}\{Z\} / x_i \}^{-1/(2D)} \quad 13$$

see Friston et al (1991) and Worsley et al (1992) for more details.

*The p-value based on n, the size of the region*

The probability of getting one or more regions of say size  $k$  or more in a given  $\text{SPM}\{Z\}$  thresholded at  $u$  (say  $\text{SPM}_u\{Z\}$ ) of volume  $S$  [ $P_S(n > k)$ ] is the same as the probability that the largest region consists of  $k$  or more voxels [ $P(n_{\max} > k)$ ]. Expressions for this probability rely on a number of previously known distributional approximations and a conjecture that  $n^{2D}$  has an exponential distribution (in the limit of high thresholds) (Nosko 1969, 1970; Adler 1981). By assuming this form for  $P(n = x)$  to be asymptotically correct we (Friston et al 1994) determined the parameters of the distribution by reference to its known moments. It was shown that:

$$P_S(n > k) = P(n_{\max} > k) = 1 - \exp(-E\{m\} \cdot e^{-k^{2D}})$$

$$\text{where } E\{m\} = \left[ (D/2 + 1) \cdot E\{m\} / S \cdot (-u) \right]^{2/D} \quad 14$$

Where  $(-u)$  is the error function (integral of the unit Gaussian distribution) evaluated at the threshold chosen  $(-u)$ . Eqn(14) gives an estimate of the probability of finding at least one region with  $k$  or more voxels in an  $\text{SPM}_u\{Z\}$ . Notice that  $u$  (the threshold) can be chosen. The optimum threshold should maximize sensitivity. In a previous paper we derived an approximate expression for the sensitivity to a 'random' signal (Friston et al 1994). We demonstrated how the sensitivity, or power, depends on an interplay between the shape of the signal and the threshold used. Our results indicated that low thresholds are more powerful for activation foci that are larger than the FWHM of the imaging device. In the sense that all signals (that are not subject to partial volume effects) arise in structures that satisfy this constraint one might anticipate

that low thresholds will be generally the more powerful. On the other hand, if the FWHM is bigger than at least some focal activations, the significance based on peak height may be more powerful. In what follows we present both characterizations

It should be born in mind that the results presented here are only good approximations for large thresholds, so it would seem prudent to keep the thresholds high enough for the p-values to remain valid. Simulations suggest that the approximations hold reasonably well for thresholds as low as 2.4 (see Friston et al 1994).

## **Applications**

This section describes the data used to illustrate the diversity of ways the expressions above can be used. The PET data were obtained from normal subjects during a word generation activation paradigm. The results of each analysis will be displayed in the same format. This format includes the design matrix, the contrast, the SPM{Z} and a table of regions that have been characterized in terms of their size and their maximal height [ $P_S(n_{\max} > k)$  and  $P_S(Z_{\max} > u)$ ].

### *The data*

The data were obtained from five subjects scanned 12 times (every eight minutes) whilst performing one of two verbal tasks. Scans were obtained with a CTI PET camera (model 953B CTI Knoxville, TN USA).  $^{15}\text{O}$  was administered intravenously as radiolabelled water infused over two minutes. Total counts per voxel during the buildup phase of radioactivity served as an estimate of regional cerebral blood flow (rCBF) (Fox and Mintun 1989). Subjects performed two tasks in alternation. One task involved repeating a letter presented aurally at one per two seconds (*word shadowing*). The other was a paced verbal fluency task, where the subjects responded with a word that began with the letter presented (*intrinsic word generation*). To facilitate intersubject pooling, the data were realigned and spatially [stereotactically] normalized (Friston et al submitted) and smoothed with an isotropic Gaussian kernel (FWHM of 16mm).

### *A single subjects analysis*

This example introduces the basic implementation and highlights an equivalence between various simple statistical tests (e.g. unpaired t-tests and testing for correlations). It deals with situations in which there is one subject scanned in many conditions or (equivalently from a mathematical perspective) many subjects scanned under the same condition. This section also covers comparing two groups of scans with paired and unpaired t tests. Finally it introduces the notion of removing confounding effects using linear regression.

In this the first and simplest example we address the effects of activations due to intrinsic word generation or, equivalently, deactivations due to word repetition (extrinsic word generation) in a single subject (one of the five studied). Following the philosophy of cognitive subtraction this is effected by subtracting the word shadowing from the verbal fluency conditions to assess the activations associated with cognitive components in word generation that are not in word shadowing (e.g. the intrinsic generation of word representations and the 'working memory' for words already produced). In this instance we have one *study*, one *subject* and 12 *conditions*. (comprising two *tasks*). The condition effects of interest are tested using a covariate of the form  $\mathbf{G}_c = [-1 \ 1 \ -1 \ \dots \ 1]^T$  i.e. -1 for word shadowing and 1 for word generation. Of course we can treat this covariate as levels in a factor and  $\mathbf{G}_c$  could be renamed  $\mathbf{G}_1$ . As stated in the theory section there is no mathematical distinction: In this instance there is no difference between regressing rCBF on a series of +1s and -1s or taking the mean difference between odd and even scans. To account for the confounding effects of global differences the global activities were considered as covariates of no interest  $\mathbf{H}_c$ . There are two covariate effects to be estimated (one interesting and the other not). The analysis can be seen as an example of (i) multiple linear regression, (ii) ANCOVA (regarding  $\mathbf{G}_c$  as  $\mathbf{G}_1$ ) or (iii) a simple correlation after partialling out a 'nuisance'

variable or confound. The important point is that these perspectives are all rather redundant because *they are all exactly the same instantiation of the same linear model*.

The design matrix  $\mathbf{G} = [\mathbf{G}_c \mid \mathbf{H}_c]$  corresponding to this analysis is seen in Figure 1 (top right). The resulting SPM{Z} (top left) is seen to identify significant actions in the left dorsolateral prefrontal cortex (including the opercular portion of Broca's area and related insula), left extrastriate areas, right cerebellum, right and left frontal pole and the precuneus. Note that the contrast is simply '1' for the covariate effect of interest and zero elsewhere  $\mathbf{c} = [1 \ 0]$ . In general the p-values based on spatial extent are less 'significant' than those based on peak height. This would be expected if the regions activated were smaller than the FWHM of the SPM [about 18 mm according to eqn(13)].

In the above analysis we could have pretended that we had studied 12 individuals in one of two states; with two *studies*, each with six *subjects* scanned in a single *condition*. This has exactly the same degrees of freedom and gives exactly the same SPM{Z} as above and (if we ignore the confounding covariate) can be seen as an unpaired t-test at each voxel. If one had scanned 6 subjects under *both* tasks there would be one *study*, six *subjects* and 2 *conditions*. The degrees of freedom in this example would be less because one would also be estimating subject-specific (block) effects and the design matrix would include a subject or block partition  $\mathbf{H}_1$  (c.f. a paired t-test). In general the degrees of freedom of the t statistic is equal to the number of scans minus the number of estimated parameters (including a constant term, that may be explicitly included in  $\mathbf{H}_1$  or made implicit in the analysis by mean correction of the data and covariates).

In the analysis of groups of subjects with scans obtained under the same condition this design (using one covariate of interest) can be useful in indentifying regionally specific associations between physiology and some behavioural or symptom score in a parametric fashion (e.g. Friston et al 1992a).

### *An activation study using intersubject averaging*

This example introduces the more general ANOVA-like layout and its extension to ANCOVA, again using covariates to remove the effects of confounding or nuisance variables. This removal reveals more clearly the underlying condition-specific effects of interest and is exactly the same as that implemented in the previous section (i.e. by linear regression). Three sorts of analyses are considered (*subtractive, parametric and factorial*). All are implemented at the level of the contrast used in comparing condition-specific effects.

In this case there is one *study*, five *subjects* and 12 *conditions*. We removed the confounding effects of global activity by designating these as covariates of no interest  $\mathbf{H}_c$ . This example is equivalent to a one way ANCOVA with a completely blocked design. There are 12 condition specific effects, five subject effects (block) and a covariate effect. The SPM{F} corresponding to this analysis is depicted in Figure 2. This SPM can be regarded as an image of the [significance of] variance introduced by the experimental design (or more exactly the sums of squares due to condition effects relative to error). The design matrix used in this example is seen in Figure 3 and is displayed by placing the partitions in the same order as in Eqn(4) i.e. [  $\mathbf{G}_1$  |  $\mathbf{G}_c$  |  $\mathbf{H}_1$  |  $\mathbf{H}_c$  ]. Here  $\mathbf{G}_1$  models the 12 conditions and  $\mathbf{G}_c$  is empty because there are no covariates of interest. Because of the constraint on the design matrix only four subject effects are estimated directly (see the corresponding design matrix  $\mathbf{H}_1$  partition).  $\mathbf{H}_c$  is seen to be a column of global activities (on the far left). The objective of further analysis, using contrasts, is to test specific hypotheses that particular differences between conditions account for this variance in physiology:.

### *A subtractive approach*

Suppose one wanted to identify significant activation foci associated with word generation as above by pooling data from all the subjects. Having estimated the 12 condition-specific effects the effect of verbal fluency *vs* word shadowing is assessed using a contrast that is 1 in all the verbal fluency conditions, -1 in the word generation

and 0 elsewhere  $\mathbf{c} = [1 \ -1 \ 1 \ -1 \ \dots \ -1 \ 0 \ 0 \ 0 \ 0 \dots]$ . The results of this analysis are presented in Figure 3 which shows the design matrix, the contrast and the resulting  $\text{SPM}\{Z\}$ . The results demonstrate significant activations in the left anterior cingulate, left dorsolateral prefrontal cortex, operculum and related insula, thalamus and left extrastriate (among others). The relationship of the left prefrontal and mediodorsal thalamic activations, to the underlying anatomy is seen in Figure 4 which is the  $\text{SPM}\{Z\}$  sectioned in three orthogonal planes and rendered on an arbitrary structural MRI scan. The extent and regional topography of the dorsolateral prefrontal activation is highlighted in Figure 5 by rendering the SPM onto the cortical surface of the same structural scan as in the previous figure.

The correspondence between this group analysis and the single subject analysis can be seen by comparing the  $\text{SPM}\{Z\}$ s in Figures 1 and 3. The most conspicuous difference is that unlike the group as a whole, the single subject fails to activate the thalamus (see below) but overall there is a remarkable similarity.

#### *A parametric approach*

In this example we tested for monotonic and linear time effects using the contrast depicted in Figure 6. The results of this analysis identify bilateral foci in the posterior temporal regions, the precuneus and the left prefrontal and anterior cingulate that show monotonic decreases in rCBF. These decreases are task-independent.

This example is trivial in its conception but is used here to introduce the notion of a parametric approach to the data. Parametric approaches avoid many of the shortcomings of 'cognitive subtraction'; and 'additive factors logic' in testing for systematic relationships between neurophysiology and sensorimotor, psychophysical, pharmacologic or cognitive parameters. These systematic relationships are not constrained to be linear or additive and may evidence very nonlinear behaviour reflecting complex interactions at a physiological or cognitive level (from a statistical perspective these interactions are linear in the parameters).



### *A factorial approach*

This example looks at regionally specific interactions, in this instance between the activation effect due to intrinsic word generation and time. The contrast used is depicted in Figure 7 and shows a typical mirror symmetry. The design matrix has not changed but we are now testing for a specific profile of condition effects. The greatest  $t$  values will obtain when the condition effects (i.e. relative activations) 'match' the contrast. As the contrast initially goes up, down, up, down and then switches to down, up, down up, this contrast will highlight those regions that de-activate early in the experiment and activate towards the end. In other words those areas that show a time-dependent augmentation of their activation. The areas implicated include left frontal operculum, insula, thalamus and superior temporal cortex. Note that these regional results suggest a true physiological 'adaptation' in the sense that it is the physiological *response* (to a task component) that shows a time-dependent change (contrast this with the task-independent changes of the previous section).

The SPM{Z} in this analysis is reported in a *descriptive* way only because no region was assessed as significant (at  $p < 0.05$ ). This means that unless one had made specific predictions about which regions were going to be involved, *no statistical inference about regional activation could be made on the basis of these results*.

This example also shows how to do a factorial experiment without explicitly modeling interaction terms in the design matrix. This approach has proved powerful in the demonstration of time-dependent adaptation during motor practice (Friston 1992b) and in psychopharmacological activation studies crossing cognitive activation and pharmacological manipulations (Grasby et al 1993). In the example above we have described a time-dependent reorganisation of physiological responses to the same task. It is tempting to call this plasticity, however the term plasticity means many things to many people and hence the term should be used carefully.

### *The adjusted responses*

Recall that adjusted data has been adjusted for uninteresting effects [see Eqn(5)]. As an example of an adjusted response we have chosen data from a voxel from the left thalamus. This region clearly shows a variable and complicated response with time-dependent changes (see previous section). The adjusted data for the analysis of all subjects for a voxel in the left mediodorsal thalamus is shown in Figure 8. The bars represent the mean adjusted condition-specific estimates of rCBF and the dots correspond to individual (adjusted) data. The regional activation from which this voxel was selected is seen on orthogonal MRI sections in Figure 4. This thalamic response is very interesting with no activation in the first 4 conditions and a profound activation in the last six. At some point during the course of this experiment the physiological changes supporting the cognitive processing elicited by verbal fluency changed markedly to involve the mediodorsal thalamus and, by reference to figure 7, augment responsiveness of the opercular cortex and insula.

#### *The single subject analyses revisited*

This section deals with situations where one wants to compare several activation studies obtained in different groups. This sort of design is especially powerful in looking at the effect of centrally active drugs, pathophysiology, clinical diagnosis etc, on condition-specific activations, where the conditions may themselves involve several treatments and the condition-specific effects may themselves be interaction terms (e.g. the adaptation above). We take the limiting case of this general situation where there is only one subject in one of two studies. This ties in neatly with the single subject analysis above and allows us to make a number of points; however it should be remembered that everything in the remainder of this section applies when the two (or more) studies include more than one subject.

In this application we want to look at the first subject in the context of the remaining four in terms of activations due to intrinsic word generation. Although we might anecdotally compare the activation profiles for the single subject (Figure 1) and for the group (Figure 3) the apparent differences may not be the most important (or indeed

significant). This section addresses the differences between the single subject's activations and the group's activation in a more formal way. Here the design matrix has two *studies*. The first study has four *subjects* and the second study is of the single *subject*. In both studies there are 12 conditions, giving effectively 24 conditions (12 in the group and 12 for the single subject). Again global activity was treated as a confounding covariate. The differences between the activations constitute an interaction:

#### *Study x condition interaction*

The difference of interest represents a *condition x study* interaction and is tested using the contrast shown in Figure 9 (again note the mirror symmetry). This contrast will highlight regions that activate more in the group than the single subject (or deactivate less). Because there are effectively 24 conditions effects (two studies with 12 conditions) the contrast is a vector with 24 elements. The SPM{Z} in Figure 9 shows that the most significant differences are in bilateral auditory and periauditory regions. These are areas that deactivate generally and more profoundly in the single subject studied. The condition effect tested here is that due to the difference between the two tasks, however we could have tested other condition specific effects (e.g. time). This analysis directly addresses how the physiological response of a single subject differs from some normative data. In other words the activation effects peculiar to this individual's functional anatomy can be directly assessed using an interaction term. The importance of this approach lies in careful characterization of single cases when applied using a lesion deficit model or in assessing inter-individual variability in functional anatomy.

## Discussion

We have described a simple framework within the context of the general linear model that allows for a diverse interrogation of functional imaging data using statistical parametric maps. The same implementation and layout can accommodate approaches that are as simple as an unpaired  $t$  test, to ANCOVA with multiple covariates. This approach partitions the design matrix at two levels (i) according to whether the effect is interesting or not and (ii) whether the effect is factor level (an indicator-type variable) or continuous (a covariate). The first distinction is fundamental and directly affects adjustment for confounding effects and the estimation of omnibus significance at each voxel. The second distinction is purely conceptual; mathematically there is no distinction between indicators and covariates.

Specific effects modelled in the design matrix are assessed using linear compounds (contrasts) of the parameter estimates (such as condition-specific mean activity or regression coefficients). The resulting statistic's distribution has the Student's  $t$  distribution under the null hypothesis and is used to make statistical inferences. Inferences about local excursions (peaks) of the  $SPM\{t\}$  (after transformation to a  $SPM\{Z\}$ ) use  $p$ -values that are estimated using distributional approximations from the theory of Gaussian fields. These  $p$ -values are based on the region's highest value and the number of voxels comprising that region.

This paper presents the operational expressions required to perform the analysis and some examples which cover most applications one might envisage. In particular we have focussed on comparing activation studies performed in different subjects. Our discussion uses a general taxonomy of activation studies that distinguishes between subtractive (categorical), parametric (dimensional) and factorial (interactions) designs. Subtractive designs are well established and powerful devices in functional mapping but are predicated on possibly untenable assumptions about the relationship between brain dynamics and the functional processes that ensue (and where these assumptions

may be tenable they are not demonstrated to be so). The main concerns with subtraction and additive factors logic can be reduced to the relationship between neural dynamics and cognitive processes. For example even if, from a functionalist perspective, a cognitive component can be added without interaction with pre-existing components the brain's implementation of these processes is almost certainly going to show profound interactions. This follows simply from the observation that neural dynamics are nonlinear (e.g. Aertsen and Preissl 1991). Indeed nearly all theoretical and computational neurobiology is based on this observation. Parametric approaches avoid many of the philosophical and physiological shortcomings of 'cognitive subtraction'; in testing for systematic relationships between neurophysiology and sensorimotor, psychophysical, pharmacologic or cognitive parameters. These systematic relationships are not constrained to be linear or additive and may show very nonlinear behaviour. The fundamental difference between subtractive and parametric approaches lies in treating a cognitive process, not as a categorical invariant, but as a dimension or attribute that can be expressed to a greater or lesser extent. It is anticipated that parametric designs of this type will find an increasing role in psychological and psychophysical activation experiments. Finally factorial experiments provide a rich way of assessing the effect of one manipulation on the effects of another. They could also be used to establish the validity of subtraction by assessing the degree of 'interaction' between cognitive processes at a physiological level. The assessment of differences in activations between two or more groups represents a question about regionally specific interactions. The limiting case of this example is where one group contains only one subject and we suggest that this is one way to proceed with single subject analyzes. We mean this in the sense that the interesting things about an individual's activation profile are how it relates to some normal profile or a profile obtained from the same subject in different situations or at a different time. These *differences in activations are interactions*.

#### *Assumptions and limitations*

The validity and scientific utility of statistical parametric mapping has been established by its diverse and expert application in many imaging centers over the past years. Given that the advantages are generally accepted we present here a critical evaluation of some of the assumptions and limitations of the approach:

*(i) Parametric assumptions*

Underlying the general linear model is an assumption that the error terms are normally distributed. Recent interest in non parametric approaches (e.g. Holmes et al - submitted) might be interpreted as a challenge to this assumption (this interpretation is not correct). There are a number of reasons for being confident that the data obtained with imaging devices (particularly PET) conform to Gaussian distributions. The image reconstruction process in PET (back projection) can be thought of in terms of convolving the underlying distribution of radiodecay events with itself many many times. The underlying distribution is approximately Poisson and by central limit theory the univariate distribution of intensity values in the back projected image will be, almost certainly, Gaussian. This argument does not however allow for non Gaussian behaviour of the physiological component in functional images (although there is no reason to suppose they are not Gaussian); however one can make a reasonable argument that the univariate behaviour of the final measurements will be Gaussian. This is because of explicit and implicit convolutions of the original distributions in the early parts of data processing [e.g. ramp and Hanning filtering in frequency space (i.e. convolving in Cartesian space) during reconstruction and Gaussian smoothing of images as a pre-processing step]. Even if the original physiological measurements were not Gaussian, after these convolutions they will be (nearly).

*(ii) Homoscedasticity*

Homoscedasticity is term referring to invariant second order behaviour (variance) of the error term over different conditions. Although generally accepted in most applications of the general linear model this constancy is not necessarily always the

case. If the error variance associated with one condition was very different from that associated with another then the statistical inferences based on a  $t$  value in the  $SPM\{t\}$  cannot be guaranteed. This is because the distributional approximations may no longer be valid. Given that very few observations (usually up to twelve for PET activation studies) are available it would be very difficult to demonstrate a significant difference in error variance (or indeed demonstrate that they were not significantly different). It is interesting to note that the criticism levelled at approaches that use an estimate of error variance pooled over voxels (namely, one cannot assume that error variance is region-independent) can be turned around and applied to statistical parametric mapping (namely one cannot assume that error variance is condition-independent). Variance estimators which accommodate condition-specific changes in error variance are the subject of some current work (Worsley - personal communication).

### *(iii) Stationarity*

In statistical parametric mapping weak stationarity is assured under the null hypothesis since the mean at any voxel is zero and the variance is unity, because the error variance is estimated separately at each voxel. However there is another aspect of stationarity that is assumed; namely a stationary multivariate behaviour, or more simply the correlations between different parts of the SPM are unchanging (for Gaussian fields homogeneous correlation structure, in addition to weak stationarity implies strict stationarity). This assumption is implicit in modelling the SPM as a stationary stochastic process (usually a Gaussian random field). It is possible that (i) the smoothness changes from region to region and (ii) long range correlations in error variance may show a regional specificity. The first nonstationary behaviour can only be demonstrated with 'local' estimates of smoothness (W above). The distributional approximations for the estimators of  $W$  are currently being investigated in order to address this issue at an empirical level (Poline - personal communication). It is expected that this form of nonstationarity will not be significant in the sense that one can usually assume the same point spread function for all parts of the original data. The

second sort of regionally specific deviation from stationarity is due to functional interactions between remote brain areas that are mediated physiologically (i.e. by functional connectivity). This sort of nonstationarity can be discounted at one level by noting that under the null hypothesis there are no systematic physiological interactions. A deeper analysis however reveals the fundamental distinction between the mass univariate approach taken by statistical parametric mapping and equivalent multivariate approaches. In multivariate approaches one would explicitly use the measured covariance structure of the error terms in making some statistical inference about the effects of a treatment on the multivariate measure (i.e. the profile of activity over the entire brain). Because these inferences depend on inverting the error variance-covariance matrix they are not applicable to voxel-based analyses of neuroimaging data (this is because the matrices are singular due to the great number of voxels relative to the number of scans). The solution is therefore to proceed on a univariate basis (treating each voxel as if it were measured in isolation) and then model the covariances among voxels using a Gaussian field model. Note also that the multivariate approach is rather weak in that it cannot be used to make regionally specific inferences.

*(iv) Low degrees of freedom*

SPMs will fail at low degrees of freedom. The reasons for this failure can be quite entertaining (see Worsley et al 1993b). This limitation is however simply the familiar caution 'statistics are only meaningful when one has enough data' restated in the context of spatially extended statistical processes. It is difficult to prescribe a lower limit for the degrees of freedom that one should use: Ideally one would like to see at least 30. However the practical limitations of PET make this number of scans prohibitive in the analysis of single subjects and one could be obliged to work with ten degrees of freedom or less. Because low degrees of freedom compromise conventional single subject analyses we consider the suggestions above concerning single subjects (and their analysis using group data) to be particularly important. It should be remembered that the lower the degrees of freedom the less a transformed  $SPM\{t\}$



conforms to a Gaussian field model. In this case of very low degrees of freedom (e.g. 10 - 20) non parametric approaches may be more appropriate (e.g. Holmes et al 1994) or reference to distributional approximations for the  $SPM\{t\}$  (as opposed to the  $SPM\{Z\}$ ) should be considered (Worsley 1994)

In conclusion we have presented a unified framework using the general linear model and the theory of Gaussian fields that facilitates the analysis of neuroimaging data using a variety of experimental layouts. These can range from categorical and factorial activation studies to parametric designs with many covariates. This generalization should increase the latitude of possible experiments available to the imaging neuroscientist.

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## References

- Adler RJ (1981) The geometry of random fields. Wiley, New York.
- Aertsen A & Preissl H (1991) Dynamics of activity and connectivity in physiological neuronal Networks in Non Linear Dynamics and Neuronal Networks Ed Schuster HG VCH publishers Inc., New York NY USA p 281-302
- Bench CJ Friston KJ Brown RG Scott LC Frackowiak RSJ and Dolan RD. The anatomy of Melancholia - focal abnormalities of cerebral blood flow in major depression Psychol Med (1992) 3:607-615
- Chatfield C and Collins AJ (1980) Introduction to multivariate analysis. Chapman and Hall, London. p189-210
- Fox PT Mintun MA Reiman EM and Raichle ME (1988) Enhanced detection of focal brain responses using intersubject averaging and distribution analysis of subtracted PET images J Cereb. Blood Flow 8:642-653
- Fox PT, Mintun MA (1989) Non-invasive functional brain mapping by change distribution analysis of averaged PET images of H<sup>15</sup>O<sub>2</sub> tissue activity J.Nucl. Med. 30:141-149
- Friston KJ Frith CD Liddle PF Dolan RJ Lammertsma AA & Frackowiak RSJ (1990) The relationship between global and local changes in PET scans J. Cereb. Blood Flow Metab. 10:458-466
- Friston, K.J Frith, C.D Liddle, P.F and Frackowiak, R.S.J. (1991) Comparing functional (PET) images: The assessment of significant change. J. Cereb. Blood Flow Metab. 11:690-699.
- Friston KJ Liddle PF Frith CD Hirsch SR and Frackowiak RSJ. (1992a) The left medial temporal region and schizophrenia: A PET Study. Brain 115:367-382
- Friston, K.J Frith, C Passingham, R.E Liddle, P and Frackowiak, RSJ (1992b) Motor Practice and neurophysiological adaptation in the cerebellum: a positron tomography study. Proc. Royal Soc. Lond. Series B. 248:223-228.

Friston, K.J, Grasby, P, Bench, C, Frith, C, Cowen, P, Little, P., Frackowiak, R.S.J and Dolan, R (1992c) Measuring the neuromodulatory effects of drugs in man with positron tomography. *Neuroscience Letters* 141:106-110.

Friston KJ, Frith CD, Liddle PF & Frackowiak RSJ (1993) Functional connectivity: The principal component analysis of large (PET) data sets. *J. Cereb. Blood Flow Metab.* 13:5-14

Friston KJ, Worsley KJ, Frackowiak RSJ, Mazziotta JC & Evans, AC (1994) Assessing the significance of focal activations using their spatial extent. *Human Brain Mapping* 1:214-220.

Friston KJ, Ashburner J, Poline JB, Frith CD, Heather JD and Frackowiak RSJ Spatial realignment and normalization of images, submitted.

Frith CD, Friston KJ, Liddle PF, and Frackowiak RSJ. (1991) Willed action and the prefrontal cortex in man. *Proc. R. Soc. Lond. B* 244:241-246.

Grafton S, Mazziotta J, Presty S, Friston KJ, Frackowiak RSJ and Phelps M (1992) Functional anatomy of human procedural learning determined with regional cerebral blood flow and PET. *J. Neuroscience* 12:2542-2548.

Grasby P, Friston KJ, Bench C, Cowen P, Frith C, Liddle P, Frackowiak, R.S.J and Dolan R (1992) Effect of the 5-HT<sub>1A</sub> partial agonist buspirone on regional cerebral blood flow in man. *Psychopharmacology* 108:380-386

Hasofer AM (1978) Upcrossings of random fields. *Suppl. Adv Appl. Prob.* 10:14-21

Holmes AP, Blair RC, Watson JDG and Ford I. Non-parametric analysis of statistical images from functional mapping experiments *J. Cereb. Blood Flow Metab.* - submitted

Lueck CJ, Zeki S, Friston KJ, Deiber NO, Cope P, Cunningham VJ, Lammertsma AA, Kennard C, and Frackowiak RSJ. The colour centre in the cerebral cortex of man. *Nature* (1989) 340:386-389.

Martin, AJ, Friston, KJ, Colebatch, JG, and Frackowiak RSJ. Decreases in regional cerebral blood flow in normal aging. *J.Cereb. Blood Flow Metab.* (1991) 694-689.

Nosko VP (1969) Local structure of Gaussian random fields in the vicinity of high level shins. *Soviet Mathematics Doklady*, 10:1481-1484

Nosko VP (1970) On shines of Gaussian random fields (in Russian) Vestnik Moscov. Univ. Ser. I Mat. Meh., 1970:18-22

Price C Wise RJS Ramsay S Friston KJ Howard D Patterson K and Frackowiak RSJ (1992) Regional response differences within the human auditory cortex when listening to words. Neurosci. Letters 146:179-182

Scheffe H (1959) The analysis of variance, Wiley, New York.

Spinks TJ Jones T Bailey DL Townsend DW Grootnook S Bloomfield PM Gilardi MC Casey ME Sipe B and Reed J (1992) Physical performance of a positron tomograph for brain imaging with retractable septa. Phys. Med. Biol. 37:1637-1655

Talairach J, Tournoux P (1988) A Co-planar stereotaxic atlas of a human brain. Thieme, Stuttgart

Townsend DW Geissbuhler A Defrise M Hoffman EJ Spinks TJ Bailey D Gilardi MC and Jones T (1992) Fully three-dimensional reconstruction for a PET camera with retractable septa IEEE Trans Med. Imaging M1-10:505-512

Winer BJ (1971) Statistical Principles in experimental design, 2nd Edition, McGraw-Hill, New York

Worsley KJ Evans AC Strother SC and Tyler JL (1991) A linear spatial correlation model with applications to positron emission tomography J. Am. Statistical Assoc. 86:55-67

Worsley KJ Evans AC Marrett S & Neelin P (1992) A three-dimensional statistical analysis for rCBF activation studies in human brain. J. Cereb. Blood. Flow Metab. 12:900-918

Worsley KJ Evans AC Marrett S and Neelin P (1993a) Detecting and estimating the regions of activation in CBF activation studies in human brain. In Qualification of brain function: Tracer kinetics and image analysis in brain PET. eds. K Uemura, N Lassen, T Jones and I Kanno Excerpta Medica, London p535-548

Worsley KJ, Evans EC, Marrett S and Neelin P (1993b) Authors reply (letter) J Cereb. Blood Flow Metab. 13:1041-1042

Worsley KJ (1994) Local Maxima and the expected Euler characteristic of excursion sets of  $\chi^2$ , F and t fields. *Advances Appl. Prob.* 26;13-42

## Legends for figures

### *Figure 1*

Results of a single subject analysis. The format of this Figure is the same as for Figures 1, 3, 6, 7 and 9:

**Top right:** Design matrix: This is an image representation of the design matrix, because elements of this matrix can take negative values the gray scale is arbitrary and has been scaled to the minimum and maximum. The form of the design matrix is the same as in the text [condition effects, covariates of interest, subject effects, covariates of no interest] = [ $\mathbf{G}_1$   $\mathbf{G}_c$   $\mathbf{H}_1$   $\mathbf{H}_c$ ] where these partitions exist. Contrast: This is the contrast or vector defining the linear compound of parameters tested ( $\mathbf{c}$ ). The contrast is displayed over the column of  $\mathbf{G}$  that corresponds to the effect[s] in question. Note that the length of the contrast is the same as the number of columns in the design matrix, which is the same as the number of parameters one is explicitly estimating. In this Figure there are only two parameters to be estimated and these are the regression coefficients for the covariate testing for the difference between verbal fluency and word shadowing and the confounding covariate of global activity.

**Top left:** SPM{Z}: This is a maximum intensity projection of the SPM{t} following transformation to the Z score. The display format is standard and provides three views of the brain from the front, the back and the right hand side. The grayscale is arbitrary and the space conforms to that described in the atlas of Talairach and Tournoux (1988).

**Lower panel:** Tabular data are presented of 'significant' regions ( $p < 0.05$  corrected the volume of the SPM{Z}). The location of the maximal voxel in each region is given with the size of the regions (n), and the peak Z score. For each region the significance is assessed in terms of  $E\{m\} > P(Z_{\max} > Z)$  using eqn(11) and  $P(n_{\max} > n)$  using eqn(14). In this figure there are eight significant regions that are described in the text. The footnote gives the volume in voxels of 2 x 2 x 4 mm and the degrees of freedom.

### *Figure 2*

SPM{F}: Maximum intensity projection of the SPM{F} computed according to eqn(9) for the design matrix in Figure 3. The design matrix was used to assess the 12 condition effects in all five subjects. Experimental variance is prominent in bitemporal, [anterior and posterior] cingulate, thalamic regions and opercular portions of Broca's area. The display format is the same as in the previous figure and the gray scale is arbitrary. All the F ratios are significant at  $p < 0.05$  (uncorrected).

### *Figure 3*

A subtractive analysis based on intersubject averaging.

The format is the same as for Figure 1. The design matrix now includes condition and subject (block) partitions and the contrast can be seen to test for differences in condition means in the verbal fluency and word shadowing conditions. The corresponding profile of activations is seen in the SPM{Z}. Significant activations are tabulated below and are described in the main text.

### *Figure 4*

The same data presented in Figure 3 but here the SPM{Z} has been sectioned in three orthogonal planes and displayed on top of an arbitrary MRI image that has been spatially normalized to the same anatomical space. This figure details the functional anatomy of the left prefrontal and mediodorsal thalamic activations and their relationship to underlying anatomy.

### *Figure 5*

The same data as in Figure 3, but here the SPM{Z} has been rendered onto the same MRI image as in the previous figure (Figure 4). This figure details further the topography of the left prefrontal activation which can be seen to involve the opercular portion of Broca's area, inferior frontal gyrus and extend almost to the frontal pole. This is the largest contiguous region of activations elicited by this comparison (see Table in Figure 3).

*Figure 6*

A parametric analysis

The format of this figure is the same as for Figure 1 and shows the results of testing for a linear monotonic time effect using a contrast of the condition effect estimates

*Figure 7*

A factorial analysis

The format of this figure is the same as for Figure 1 and shows the results of testing for an interaction between activations due to verbal fluency and time. The contrast used detects regions whose response to verbal fluency increases with time. No region can be considered significant because the largest region would have been obtained on 8.5% of occasions by chance (see lower panel). The results of this analysis can only be reported descriptively and no statistical inference can be made about the regionally specific effects.

*Figure 8*

Adjusted regional activity.

Using the results displayed in the previous figure a voxel in the mediodorsal thalamus was selected and the adjusted activity was plotted for each of the 12 conditions. The bars represent mean condition-specific estimates and the dots represent individual data adjusted for confounding effects. Note the time-dependent augmentation of activations due to verbal fluency.

*Figure 9*

An interaction analysis comparing a single subject and the remaining subjects.

The design matrix now has 24 condition effects (12 for the group and 12 for the single subject). The contrast is testing for a failure to activate (or a greater de-activation) in the



single subject with respect to the group. The bitemporal areas deactivate more profoundly than one would have predicted on the basis of the group data.