

DEPARTMENT OF STATISTICS

University of Wisconsin

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## Statistical Model Building, Machine Learning, and the Ah-Ha Moment

Grace Wahba<sup>1</sup>

Department of Statistics, Department of Computer Sciences  
and Department of Biostatistics and Medical Informatics  
University of Wisconsin, Madison

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## Preamble

As a recipient of the Elizabeth Scott Award, I received the following invitation from David Scott and COPSS:

“The Committee of Presidents of Statistical Societies (COPSS) will celebrate its 50th Anniversary in 2013. As part of its celebration, COPSS intends to publish a book with contributions from the past recipients of its four awards, namely the Fisher Lecture Award, the President’s Award, the Elizabeth Scott Award, and the FN David Award. The theme of the book is Past, Present and Future of Statistical Science. As a past winner of one of the COPSS awards, we would like to invite you to contribute to this book... I will be working with you personally on your contribution ... Specifically, we are seeking contributions along (one of) the following lines:

1. Statistical Research: Your reflection on the past, present or future of a statistical research area of your choice. You are free to decide what you would like to focus on, i.e., past, present or future, or possibly all three.
2. Statistical Education: Your reflection/view on statistical education in some areas, e.g., BIG DATA, interdisciplinary research, graduate and undergraduate curriculum, promoting statistical education and research in developing countries, or statistical educational outreach.
3. Statistical Career: Your reflection on your own career, lessons and experience you have learned, and advice you would like to provide to young statisticians if sought.
4. A blend of the above three topics. ”

(signed by David Scott representing COPSS).

I have chosen to focus on Item 3, reflecting on some particular “fun” pieces of my own statistical career, sprinkled with advice, and augmented with a few assorted remarks. Following is my contribution.

# Chapter 1

## Statistical Model Building, Machine Learning, and the Ah-Ha Moment

[March 26, 2013 Chapter by Grace Wahba for the Committee of Presidents of Statistical Societies (COPSS) 50th anniversary volume]

Highly selected “Ah-Ha” moments from the beginning to the present of my research career are recalled - these are moments when the main idea just popped up instantaneously, sparking sequences of future research activity- almost all of these moments crucially involved discussions/interactions with others. Along with a description of these moments we give unsought advice to young statisticians. We conclude with remarks on issues relating to statistical model building/machine learning in the context of human subjects data.

### 1.1 Introduction-Manny Parzen and RKHS

Many of the “Ah-Ha” moments below involve Reproducing Kernel Hilbert Spaces (RKHS) so we begin there. My introduction to RKHS came while attending a class given by Manny Parzen on the lawn in front of the old Sequoia Hall at Stanford around 1963. See [24].

For many years RKHS [1, 33] were a little niche corner of research which suddenly became popular when their relation to Support Vector Machines (SVMs) became clear- more on that later. To understand most of the Ah-Ha moments it may help to know a few facts about RKHS which we now give.

An RKHS is a Hilbert space  $\mathcal{H}$  where all of the evaluation functionals are bounded linear functionals. What this means is the following: Let the domain of  $\mathcal{H}$  be  $\mathcal{T}$ , and the inner

product  $\langle \cdot, \cdot \rangle$ . Then, for each  $t \in \mathcal{T}$  there exists an element, call it  $K_t$  in  $\mathcal{H}$ , with the property  $f(t) = \langle f, K_t \rangle$  for all  $f$  in  $\mathcal{H}$ .  $K_t$  is known as the representer of evaluation at  $t$ . Let  $K(s, t) = \langle K_s, K_t \rangle$ ; this is clearly a positive definite function on  $\mathcal{T} \otimes \mathcal{T}$ . By the Moore-Aronszajn theorem, every RKHS is associated with a unique positive definite function, as we have just seen. Conversely, given a positive definite function, there exists a unique RKHS (which can be constructed from linear combinations of the  $K_t, t \in \mathcal{T}$  and their limits). Given  $K(s, t)$  we denote the associated RKHS as  $\mathcal{H}_K$ . Observe that nothing has been assumed concerning the domain  $\mathcal{T}$ . A second role of positive definite functions is as the covariance of a zero mean Gaussian stochastic process on  $\mathcal{T}$ . In a third role that we will come across later - let  $O_i, i = 1, 2, \dots, n$  be a set of  $n$  abstract objects. An  $n \times n$  positive definite matrix can be used to assign pairwise squared Euclidean distances  $d_{ij}$  between  $O_i$  and  $O_j$  by  $d_{ij} = K(i, i) + K(j, j) - 2(K(i, j))$ . In Sections 1.1.1-1.1.9 we go through some Ah-Ha moments involving RKHS, positive definite functions and pairwise distances/dissimilarities. Section 1.2 discusses sparse models and the LASSO. Section 1.3 has some remarks involving complex interacting attributes, the “Nature-Nurture” debate, Personalized Medicine, Human subjects privacy and scientific literacy, and we end with conclusions in Section 1.4.

I end this section by noting that Manny Parzen was my thesis advisor, and Ingram Olkin was on my committee. My main advice to young statisticians is: Choose your advisor and committee carefully, and be as lucky as I was.

### 1.1.1 George Kimeldorf and the Representer Theorem

Back around 1970 George Kimeldorf and I both got to spend a lot of time at the Math Research Center at the University of Wisconsin-Madison (the one that later got blown up as part of the anti-Vietnam-war movement). At that time it was a hothouse of spline work, headed by Iso Schoenberg, Carl deBoor, Larry Schumaker and others, and we thought that smoothing splines would be of interest to statisticians. The smoothing spline of order  $m$  was the solution to: find  $f$  in the space of functions with square integral  $m$ th derivative to minimize

$$\sum_{i=1}^n (y_i - f(t_i))^2 + \lambda \int_0^1 (f^{(m)}(t))^2 dt, \quad t_i, i = 1, \dots, n \in [0, 1]. \quad (1.1)$$

Professor Schoenberg many years ago had characterized the solution to this problem as a

piecewise polynomial of degree  $2m - 1$  satisfying some boundary and continuity conditions.

Our Ah-Ha moment came when we observed that the space of functions with square integrable  $m$ th derivative on  $[0, 1]$  was an RKHS with *seminorm*  $\|Pf\|$  defined by  $\|Pf\|^2 = \int_0^1 (f^{(m)}(t))^2 dt$  and with an associated  $K(s, t)$  that we could figure out. (A seminorm is exactly like a norm except that it has a non-trivial null space, here the null space of this seminorm is the span of the polynomials of degree  $m - 1$  or less.) Then by replacing  $f(t)$  by  $\langle K_t, f \rangle$  it was not hard to show by a very simple geometric argument that the minimizer of (1.1) was in the span of the  $K_t, t = t_1, \dots, t_n$  and a basis for the null space of the seminorm. But furthermore, the very same geometrical argument could be used to solve the more general problem: find  $f \in \mathcal{H}_K$ , an RKHS, to minimize

$$\sum_{i=1}^n C(y_i, L_i f) + \lambda \|Pf\|_K^2 \tag{1.2}$$

where  $C(y_i, L_i f)$  is convex in  $L_i f$ , with  $L_i$  a bounded linear functional in  $\mathcal{H}_K$  and  $\|Pf\|_K^2$  a seminorm in  $\mathcal{H}_K$ . A bounded linear functional is a linear functional with a representer in  $\mathcal{H}_K$ , that is, there exists  $\eta_i \in \mathcal{H}_K$  such that  $L_i f = \langle \eta_i, f \rangle$  for all  $f \in \mathcal{H}_K$ . The minimizer of (1.2) is in the span of the representers  $\eta_i$  and a basis for the null space of the seminorm. That is known as the representer theorem, which turned out to be a key to fitting (mostly continuous) functions in an infinite dimensional space, given a finite number of pieces of information. There were two things I remember about our excitement over the result: One of us, I'm pretty sure it was George, thought the result was too trivial and not worthwhile to submit, but submit it we did and it was accepted [12] without a single complaint, within three weeks. I have never since then had another paper accepted by a refereed journal within three weeks and without a single complaint. Advice: If you think it is worthwhile, submit it.

### 1.1.2 Svante Wold and Leaving-Out-One

Following Kimeldorf and Wahba, it was clear that for practical use, a method was needed to choose the smoothing or tuning parameter  $\lambda$  in (1.1). The natural goal was to minimize the mean square error over the function  $f$ , for which its values at the data points would be the proxy. In 1974 Svante Wold visited Madison, and we got to mulling over how to choose  $\lambda$ . It so happened that Mervyn Stone gave a colloquium talk in Madison, and Svante and I were sitting next to each other as Mervyn described using leaving-out-one to decide

on the degree of a polynomial to be used in least squares regression. We looked at each other at that very minute and simultaneously said something, I think it was “Ah-Ha”, but possibly “Eureka”. In those days computer time was \$600/hour and Svante wrote a computer program to demonstrate that leaving-out-one did a good job. It took the entire Statistics department’s computer money for an entire month to get the results in [35]. Advice: Go to the colloquia, sit next to your research pals.

### 1.1.3 Peter Craven, Gene Golub and Michael Heath and GCV

After much struggle to prove some optimality properties of leaving-out-one, it became clear that it couldn’t be done in general. Considering the data model  $y = f + \epsilon$ , where  $y = (y_1, \dots, y_n)^T$ ,  $f = (f(t_1), \dots, f(t_n))^T$  and  $\epsilon = (\epsilon_1, \dots, \epsilon_n)^T$  is a zero mean i.i.d. Gaussian random  $n$ -vector then the information in the data is unchanged by multiplying left and right hand side by an orthogonal matrix, since  $\Gamma\epsilon$  with  $\Gamma$  orthogonal is still white Gaussian noise. But leaving-out-one can give you a different answer. To explain, we define the *influence matrix*: Let  $f_\lambda$  be the minimizer of (1.1) when  $C$  is sum of squares. The influence matrix relates the data to the prediction of the data,  $f_\lambda = A(\lambda)y$ , where  $f_\lambda = (f_\lambda(t_1), \dots, f_\lambda(t_n))$ . A heuristic argument fell out of the blue, probably in an attempt to explain some things to students, that rotating the data so that the influence matrix was constant down the diagonal, was the trick. The result was that instead of leaving-out-one, one should minimize the GCV function  $V(\lambda) = \frac{\sum_{i=1}^n (y_i - f(t_i))^2}{(\text{trace}(I - A(\lambda)))^2}$  [4, 8]. I was on Sabbatical at Oxford in 1975 and Gene was at ETH visiting Peter Huber, who had a beautiful house in Klosters, the fabled ski resort. Peter invited Gene and me up for the weekend, and Gene just wrote out the algorithm in [8] on the train from Zurich to Klosters while I snuck glances at the spectacular scenery. Gene was a much loved mentor to lots of people. He was born on February 29, 1932 and died in November of 2007. On February 29 and March 1, 2008 his many friends held memorial birthday services at Stanford and 30 other locations around the world. Ker-Chau Li [17, 18, 19] and others later proved optimality properties of the GCV and popular codes in R will compute splines and other fits using GCV to estimate  $\lambda$  and other important tuning parameters. Advice: Pay attention to important tuning parameters since the results can be very sensitive to them. Advice: Appreciate mentors like Gene if you are lucky enough to have such great mentors.

### 1.1.4 Didier Girard, Mike Hutchinson, Randomized Trace and the Degrees of Freedom for Signal

Brute force calculation of the trace of the influence matrix  $A(\lambda)$  can be daunting to compute directly for large  $n$ . Let  $f_\lambda^y$  be the minimizer of (1.1) with the data vector  $y$  and let  $f_\lambda^{y+\delta}$  be the minimizer of (1.1) given the perturbed data  $y + \delta$ . Note that  $\delta^T(f_\lambda^y - f_\lambda^{y+\delta}) = \delta^T A(\lambda)(y + \delta) - A(\lambda)(y) = \sum_{i,j=1}^n \delta_i \delta_j a_{ij}$ , where  $\delta_i$  and  $a_{ij}$  are the components of  $\delta$  and  $A(\lambda)$  respectively. If the perturbations are i.i.d. with variance 1, then the expected value of this sum is an estimate of trace  $A(\lambda)$ . This simple idea was proposed simultaneously in [6, 11], with further theory in [7]. It was a big Ah-Ha when I saw these papers because further applications were immediate. In [32], p 139, I defined the trace of  $A(\lambda)$  as the “Equivalent degrees of freedom for signal”, by analogy with linear least squares regression with  $p < n$  where the influence matrix is a rank  $p$  projection operator. The degrees of freedom for signal is an important concept in linear and nonlinear nonparametric regression, and it was a mistake to hide it inconspicuously in [32]. Brad Efron later [5] gave an alternative definition of degrees of freedom for signal. The definition in [32] depends only on the data, Efron’s is essentially an expected value. Note that in the model (1.1),  $\text{trace } A(\lambda) = \sum_{i=1}^n \frac{\partial \hat{y}_i}{\partial y_i}$ , here  $\hat{y}_i$  is the predicted value of  $y_i$ . This definition can reasonably be applied to a problem with a nonlinear forward operator (that is, that maps data onto the predicted data) when the derivatives exist, and the randomized trace method is reasonable for estimating the degrees of freedom for signal, although care should be taken concerning the size of  $\delta$ . Even when the derivatives don’t exist the randomized trace can be a reasonable way of getting at the degrees of freedom for signal, see for example [34].

### 1.1.5 Yuedong Wang, Chong Gu and Smoothing Spline ANOVA

Sometime in the late 80’s or early 90’s I heard Graham Wilkinson expound on ANOVA (Analysis of Variance), where data was given on a regular  $d$ -dimensional grid:  $y_{ijk}, t_{ijk}, i = 1, \dots, I, j = 1, \dots, J, k = 1, \dots, K$ , for  $d = 3$  and so forth. That is, the domain is the Cartesian product of several one-dimensional grids. Graham was expounding on how fitting a model from observations on such a domain could be described as set of orthogonal projections based on averaging operators, resulting in main effects, two factor interactions, etc. “Ah-Ha” I thought, we should be able to do exactly same thing and more where the domain is the Cartesian product  $\mathcal{T} = \mathcal{T}_1 \otimes \mathcal{T}_2 \otimes \dots \otimes \mathcal{T}_d$  of  $d$  arbitrary domains. We want to fit functions

on  $\mathcal{T}$ , with main effects (functions of one variable), two factor interactions (functions of two variables), and possibly more terms given scattered observations, and we just need to define averaging operators for each  $\mathcal{T}_\alpha$ . Brainstorming with Yuedong Wang and Chong Gu fleshed out the results. Let  $\mathcal{H}^\alpha, \alpha = 1, \dots, d$  be  $d$  RKHSs with domains  $\mathcal{T}_\alpha$ , each  $\mathcal{H}^\alpha$  containing the constant functions.  $\mathcal{H} = \mathcal{H}^1 \otimes \dots \otimes \mathcal{H}^d$  is an RKHS with domain  $\mathcal{T}$ . For each  $\alpha = 1, \dots, d$ , construct a probability measure  $d\mu_\alpha$  on  $\mathcal{T}_\alpha$ , with the property that the symbol  $(\mathcal{E}_\alpha f)(t)$ , the averaging operator, defined by

$$(\mathcal{E}_\alpha f)(t) = \int_{\mathcal{T}(\alpha)} f(t_1, \dots, t_d) d\mu_\alpha(t_\alpha),$$

is well defined and finite for every  $f \in \mathcal{H}$  and  $t \in \mathcal{T}$ . Consider the decomposition of the identity operator:

$$I = \prod_{\alpha} (\mathcal{E}_\alpha + (I - \mathcal{E}_\alpha)) \quad (1.3)$$

$$= \prod_{\alpha} \mathcal{E}_\alpha + \sum_{\alpha} (I - \mathcal{E}_\alpha) \prod_{\beta \neq \alpha} \mathcal{E}_\beta + \sum_{\alpha < \beta} (I - \mathcal{E}_\alpha)(I - \mathcal{E}_\beta) \prod_{\gamma \neq \alpha, \beta} \mathcal{E}_\gamma + \dots + \prod_{\alpha} (I - \mathcal{E}_\alpha). \quad (1.4)$$

This decomposition of the identity then always generates a unique (ANOVA-like) decomposition of  $f \in \mathcal{H}$  of the form

$$f(t) = \mu + \sum_{\alpha} f_{\alpha}(t_{\alpha}) + \sum_{\alpha < \beta} f_{\alpha\beta}(t_{\alpha}, t_{\beta}) + \sum_{\alpha < \beta < \gamma} f_{\alpha\beta\gamma}(t_{\alpha}, t_{\beta}, t_{\gamma}) + \dots \quad (1.5)$$

where the expansion is unique and (usually) truncated in some manner in practice. Here  $\mu = (\prod_{\alpha} \mathcal{E}_{\alpha})f$ ,  $f_{\alpha} = ((I - \mathcal{E}_{\alpha}) \prod_{\beta \neq \alpha} \mathcal{E}_{\beta})f$ ,  $f_{\alpha\beta} = ((I - \mathcal{E}_{\alpha})(I - \mathcal{E}_{\beta}) \prod_{\gamma \neq \alpha, \beta} \mathcal{E}_{\gamma})f$ , etc, are the mean, main effects, two factor interactions, etc. The result is usually called an SS ANOVA model, although the components are not limited to splines. For details on how to fit the terms see [9, 10, 33, 36] and the `assist` and `gss` codes in R. Note that *nothing* has been said about  $\mathcal{T}$  and very little regarding  $\mathcal{H}^\alpha$ , other than that the constant functions are in each of the constituent spaces and averaging operators can be defined.

### 1.1.6 Vladimir Vapnik, the Mystery Caller and the SVM

The AMS-IMS-SIAM Joint Summer Research Conference on Adaptive Selection of Models and Statistical Procedures was held on the campus of Mount Holyoke College in South Hadley, Massachusetts on Sunday, June 23 through Thursday, June 27, 1996. On one of those fine days a session met on a grassy lawn of Mount Holyoke College, when Vladimir



Vapnik and I were both invited speakers. I talked first, and noted how the solution to the optimization problem (1.2) led to a function involving the span of the representers. Vladimir spoke next, describing the support vector machine (SVM), a well known and highly successful method for classification, describing something he called the “kernel trick”. He exhibited an SVM that was fitted in the span of representers in an RKHS. We will explain the SVM in a moment, but the original SVM, as proposed by Vapnik and coworkers [30] was derived from an argument nothing like what I am about to give. Somewhere during Vladimir’s talk, an unknown voice towards the back of the audience called out “That looks like Grace Wahba’s stuff.” It looked obvious that the SVM as proposed by Vapnik with the “kernel trick”, could be obtained as the the solution to the optimization problem of (1.2) with  $C(y_i, L_i f)$  replaced by the so called hinge function,  $(1 - y_i f(t_i))_+$ , where  $(\tau)_+ = \tau$  if  $\tau > 0$  and 0 otherwise. Each data point is coded as  $\pm 1$  according as it came from the “plus” class or the “minus” class. For technical reasons the null space of the penalty function consists at most of the constant functions. Thus it follows that the solution is in the span of the representers  $K_{t_i}$  from the chosen RKHS plus possibly a constant function. Yi Lin and coworkers [20, 21] showed that the SVM was *estimating the sign of the log odds ratio*, just what is needed for two class classification. The SVM may be compared to the case where one desires to estimate the *probability* that an object is in the plus class. If one begins with the penalized log likelihood of the Bernoulli distribution and codes the data as  $\pm 1$  instead of the usual coding as 0 or 1, then we have the same optimization problem with  $C(y_i, f(t_i)) = \log(1 + e^{-y_i f(t_i)})$  instead of  $(1 - y_i f(t_i))_+$  with solution in the same finite dimensional space, but it is estimating the log odds-ratio, as opposed to the *sign* of the log odds ratio. It was actually a big deal that the SVM could be directly compared with penalized likelihood with Bernoulli data, and it provided a pathway for statisticians and computer scientists to breach a major divide between them on the subject of classification, and to understand each others’ work.

For many years before the Hadley meeting, Olvi Mangasarian and I would talk about what we were doing in classification, neither of us having any understanding of what the other was doing. Olvi complained that the statisticians dismissed his work, but it turned out that what he was doing was related to the SVM and hence perfectly legitimate not to mention interesting, from a classical statistical point of view. Statisticians and computer scientists have been on the same page on classification ever since.

It is curious to note that several patents have been awarded for the SVM. One of the early ones, issued on July 15, 1997 is “5649068 Pattern recognition system using support

vectors”

I’m guessing that the unknown volunteer was David Donoho.

Advice: Keep your eyes open to synergies between apparently disparate fields.

### 1.1.7 Yoonkyung Lee, Yi Lin and the Multicategory SVM

For classification, when one has  $k > 2$  classes it is always possible to apply an SVM to compare membership in one class versus the rest of the  $k$  classes, running through the algorithm  $k$  times. In the early 2000s there were many papers on one-vs-rest, and designs for subsets vs other subsets, but it is possible to generate examples where essentially no observations will be identified as being in certain classes. Since one-vs-rest could fail in certain circumstances it was something of an open question how to do multicategory SVMs in one optimization problem that did not have this problem. Yi Lin, Yoonkyung Lee and I were sitting around shooting the breeze and one of us said “how about a sum-to-zero constraint?” and the other two said “Ah-Ha”!, or, at least that’s the way I remember it. The idea is to code the labels as  $k$ -vectors, with a 1 in the  $r$ th position and  $-1/(k-1)$  in the  $k-1$  other positions for a training sample in class  $r$ . Thus, each observation vector satisfies the sum-to-zero constraint. The idea was to fit a vector of functions satisfying the same sum-to-zero constraint. The multicategory SVM fit estimates  $f(t) = (f_1(t), \dots, f_k(t))$ ,  $t \in \mathcal{T}$  subject to the sum-to-zero constraint everywhere and the classification for a subject with attribute vector  $t$  is just the index of the largest component of the estimate of  $f(t)$ . See [14, 15, 16]. Advice: Shooting the breeze is good.

### 1.1.8 Fan Lu, Steve Wright, Sunduz Keles, Hector Corrada Bravo, and Dissimilarity Information

We return to the alternative role of positive definite functions as a way to encode pairwise distance observations. Suppose we are examining  $n$  objects  $O_i$ ,  $i = 1, \dots, n$  and are given some noisy or crude observations on their pairwise distances/dissimilarities, which may not satisfy the triangle inequality. The goal is to embed these objects in a Euclidean space in such a way as to respect the pairwise dissimilarities as much as possible. Positive definite matrices encode pairwise squared distances  $d_{ij}$  between  $O_i$  and  $O_j$  as

$$d_{ij}(K) = K(i, i) + K(j, j) - 2K(i, j), \quad (1.6)$$

and, given a non-negative definite matrix of rank  $d \leq n$ , can be used to embed the  $n$  objects in a Euclidean space of dimension  $d$ , centered at 0 and unique up to rotations. We seek a  $K$  which respects the dissimilarity information  $d_{ij}^{obs}$  while constraining the complexity of  $K$  by:

$$\min_{K \in S_n} \sum |d_{ij}^{obs} - d_{ij}(K)| + \lambda \text{trace} K \quad (1.7)$$

where  $S_n$  is the convex cone of symmetric positive definite matrices. I looked at this problem for an inordinate amount of time seeking an analytic solution but after a conversation with Vishy (S. V. N. Vishwanathan) at a meeting in Rotterdam in August of 2003 I realized it wasn't going to happen. The Ah-Ha moment came about when I showed the problem to Steve Wright, who right off said it could be solved numerically using recently developed convex cone software. The result so far is [3, 22]. In [22] the objects are protein sequences and the pairwise distances are BLAST scores. The fitted kernel  $K$  had three eigenvalues that contained about 95% of the trace, so we reduced  $K$  to a rank 3 matrix by truncating the smaller eigenvalues. Clusters of four different kinds of proteins were readily separated visually in three-d plots; see [22] for the details. In [3] the objects are persons in pedigrees in a demographic study and the distances are based on Malecot's kinship coefficient, which defines a pedigree dissimilarity measure. The resulting kernel became part of an SS ANOVA model with other attributes of persons, and the model estimates a risk related to an eye disease. Advice: Find computer scientist friends.

### 1.1.9 Gabor Szekely, Maria Rizzo, Jing Kong and Distance Correlation

The last Ah-Ha experience that we report is similar to that involving the randomized trace estimate of Section 1.1.4, that is, the Ah-Ha moment came about upon realizing that a particular recent result was very relevant to what we were doing. In this case Jing Kong brought to my attention the important paper of Gabor Szekely and Maria Rizzo [27]. Briefly, this paper considers the joint distribution of two random vectors,  $X$  and  $Y$ , say, and provides a test, called distance correlation that it factors so that the two random vectors are independent. Starting with  $n$  observations from the joint distribution, let  $\{A_{ij}\}$  be the collection of double-centered pairwise distances among the  $\binom{n}{2}$   $X$  components, and similarly for  $\{B_{ij}\}$ . The statistic, called distance correlation, is the analogue of the usual sample correlation between the  $A$ 's and  $B$ 's. The special property of the test is that it is justified for  $X$  and  $Y$  in

Euclidean  $p$  and  $q$  space for arbitrary  $p$  and  $q$  with no further distributional assumptions. In a demographic study involving pedigrees [13], we observed that pairwise distance in death age between close relatives was less than that of unrelated age cohorts. A mortality risk score for four lifestyle factors and another score for a group of diseases was developed via SS ANOVA modeling, and significant distance correlation was found between death ages, lifestyle factors and family relationships, raising more questions than it answers regarding the "Nature-Nurture" debate (relative role of genetics and other attributes).

We take this opportunity to make a few important remarks about pairwise distances/dissimilarities, primarily how one measures them can be important, and getting the "right" dissimilarity can be 90% of the problem. We remark that family relationships in [13] were based on a monotone function of Malecot's kinship coefficient that was different from the monotone function in [3]. Here it was chosen to fit in with the different way the distances were used. In (1.7), the pairwise dissimilarities can be noisy, scattered, incomplete and could include subjective distances like "very close, close.." etc. not even satisfying the triangle inequality. So there is substantial flexibility in choosing the dissimilarity measure with respect to the particular scientific context of the problem. In [13] the pairwise distances need to be a complete set, and be Euclidean (with some specific metric exceptions). There is still substantial choice in choosing the definition of distance, since any linear transformation of a Euclidean coordinate system defines a Euclidean distance measure. Advice: Think about how you measure distance or dissimilarity in any problem involving pairwise relationships, it can be important.

## 1.2 Regularization Methods, RKHS and Sparse Models

The optimization problems in RKHS are a rich subclass of what can be called regularization methods, which solve an optimization problem which trades fit to the data versus complexity or constraints on the solution. My first encounter with the term "regularization" was [29] in the context of finding numerical solutions to integral equations. There the  $L_i$  of (1.2) were noisy integrals of an unknown function one wishes to reconstruct, but the observations only contained a limited amount of information regarding the unknown function. The basic and possibly revolutionary idea at the time was to find a solution which involves fit to the data

while constraining the solution by what amounted to an RKHS seminorm,  $(\int (f''(t))^2 dt)$  standing in for the missing information by an assumption that the solution was “smooth” [23, 31]. Where once RKHS were a niche subject, they are now a major component of the statistical model building/machine learning literature.

However, RKHS do not generally provide sparse models, that is, models where a large number of coefficients are being estimated but only a small but unknown number are believed to be non-zero. Many problems in the “Big Data” paradigm are believed to have, or want to have sparse solutions, for example, genetic data vectors that may have many thousands of components and a modest number of subjects, as in a case-control study. The most popular method for ensuring sparsity is probably the LASSO [2, 28]. Here a very large dictionary of basis functions  $(B_j(t), j = 1, 2, \dots)$  is given and the unknown function is estimated as  $f(t) = \sum_j \beta_j B_j(t)$  with the penalty functional  $\lambda \sum_j |\beta_j|$  replacing an RKHS square norm. This will induce many zeroes in the  $\beta_j$ , depending, among other things on the size of  $\lambda$ . Since then, researchers have commented that there is a “zoo” of proposed variants of sparsity-inducing penalties, many involving assumptions on structures in the data; one popular example is [38]. Other recent models involve mixtures of RKHS and sparsity-inducing penalty functionals. One of our contribution to this “zoo” deals with the situation where the data vectors amount to very large “bar codes”, and it is desired to find patterns in the bar codes relevant to some outcome. An innovative algorithm which deals with a humongous number of interacting patterns assuming that only a small number of coefficients are non-zero is given in [25, 26, 37].

As is easy to see here and in the statistical literature, the statistical modeler has overwhelming choices in modeling tools, with many public codes available in the software repository R and elsewhere. In practice these choices must be made with a serious understanding of the science and the issues motivating the data collection. Good collaborations with subject matter researchers can lead to the opportunity to participate in real contributions to the science. Advice: Learn absolutely as much as you can about the subject matter of the data that you contemplate analyzing. When you use “black boxes” be sure you know what is inside them.

### **1.3 Remarks on the Nature-Nurture Debate, Personalized Medicine and Scientific Literacy**

We and many other researchers have been developing methods for combining scattered, noisy, incomplete, highly heterogeneous information from multiple sources with interacting variables to predict, classify, and determine patterns of attributes relevant to a response, or more generally multiple correlated responses.

Demographic studies, clinical trials, and ad hoc observational studies based on electronic medical records, which have familial [3, 13], clinical, genetic, lifestyle, treatment and other attributes can be a rich source of information regarding the Nature-Nurture debate, as well informing Personalized Medicine, two popular areas reflecting much activity. As large medical systems put their records in electronic form interesting problems arise as to how to deal with such unstructured data, to relate subject attributes to outcomes of interest. No doubt a gold mine of information is there, particularly with respect to how the various attributes interact. The statistical modeling/machine learning community continues to create and improve tools to deal with this data flood, eager to develop better and more efficient modeling methods, and regularization and dissimilarity methods will no doubt continue to play an important role in numerous areas of scientific endeavor. With regard to human subjects studies, a limitation is the problem of patient confidentiality - the more attribute information available to explore for its relevance, the trickier the privacy issues, to the extent that de-identified data can actually be identified. It is important, however, that statisticians be involved from the very start in the design of human subjects studies.

With health related research, the US citizenry has some appreciation of scientific results that can lead to better health outcomes. On the other hand any scientist who reads the newspapers or follows present day US politics is painfully aware that a non-trivial portion of voters and the officials they elect have little or no understanding of the scientific method. Statisticians need to participate in the promotion of increased scientific literacy in our educational establishment at all levels.

### **1.4 Conclusion**

In response to the invitation from COPSS to contribute to their 50th Anniversary Celebration, I have taken a tour of some exciting moments in my career, involving RKHS and reg-

ularization methods, pairwise dissimilarities and distances, and LASSO models, dispensing un-asked for advice to new researchers along the way. I have made a few remarks concerning the richness of models based on RKHS, as well as models involving sparsity-inducing penalties with some remarks involving the Nature-Nurture Debate and Personalized Medicine. I end this contribution with thanks to my many coauthors-identified here or not, and to my terrific present and former students. Advice: Treasure your collaborators! Have great students!

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