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(Sociedad Española de Biometría)

Albacete, 5 y 6 de septiembre de 2019
Escuela Politécnica Superior de Albacete
(Edificio Infante Don Juan Manuel)
IV Jornadas Científicas de Estudiantes de la SEB

Agrupación Politécnica de Albacete

Campus de Albacete de la Universidad de Castilla-La Mancha
Albacete, España
5 y 6 de septiembre de 2019
http://www.biometricsociety.net/iv-jseb/
Welcome!

Welcome to IV JJSEB19! This event is already the fourth JJSEB congress. After successful congress in Valencia, Barcelona and Bilbao, JJSEB19 is held at the Albacete campus of the University of Castilla-La Mancha (UCLM). Edition after edition, the JJSEB has become in a meeting in which the role of young researchers in scientific spaces is promoted strongly.

Organizing this kind of event has become in one of the main activities of the Sociedad Española de Biometría (SEB), which trusts in its young members the organization of it. JJSEB promotes scientific dissemination at early stages of the scientific career and also offers students a relaxed atmosphere to present their work. Thanks to the economic support of the SEB and the BIOSTATNET network, JJSEB19 is also able to offer several stipends for scholars to help economically.

We would like to thank the Local Committee at the UCLM for the excellent local organization. We would also like to thank Virgilio Gómez-Rubio, the instructor of the specific training course as well as the Super-Women of the round table, María Durbán, Irantzu Barrio, Anabel Forte and Marta Bofill for contributing in the creation of a great program. Finally, a big thank to all of you for coming, for your support, and for helping us to make JJSEB19 a success. We wish us all a great conference! Enjoy JJSEB19 and Albacete!

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Training Course

Title: Introduction to spatial mapping with R

Instructor: Virgilio Gómez-Rubio (Universidad de Castilla-La Mancha)

Abstract:
This short course will focus on recent developments for spatial mapping and data analysis in R. Currently, R provides a myriad of R packages to handle spatial data, which includes data input/output, visualization and spatial data analysis. The course will provide an introduction to several packages for handling spatial data and making (interactive) maps. Next, the analysis of lattice and geostatistical data will be discussed using relevant examples. All course materials are available from https://github.com/becarioprecario/curso_IVJJSEB.

References:
Speakers
Development of statistical methodologies to work with ELISpot assays

Yovaninna Alarcón Soto\textsuperscript{1}, Klaus Langohr\textsuperscript{1} and Guadalupe Gómez Melis\textsuperscript{1}

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The main goal of this work is to provide statistical methodologies to analyze Enzyme-Linked Immunosorbent Spot (ELISpot) assays data. The ELISpot assay is an immunoassay that measures the frequency of cytokine-secreting cells at the single-cell level. It is carried out in a 96-well plate, and an automated ELISpot reader is used for analysis. The assay is therefore easy to perform and allows rapid analysis of a large number of samples. While ELISpot assays allows one to directly visualize and count extremely low frequencies of cytokine secreting T cells amongst millions of non-secreting bystander cells, the interpretation of ELISpot data can become ambiguous when (a) spot numbers in antigen-containing wells are low, (b) spot counts in negative control wells are elevated, and particularly (c) when both of the above occur simultaneously. Thus, the primary task, before any statistical analysis, must be the optimization of the basic assay parameters. This includes low background signal in the negative control wells, and the maximal number of antigen-induced spots in test wells, i.e., the signal to noise ratio is maximized (Dittrich and Lehmann, 2012). Our main variables of interest are spot count and spot size. Spot count corresponds to cytokine-producing cell numbers. Due to diffusion properties, a true spot has a densely coloured center which fades toward the edges; the size or color intensity of the spots is determined by the amount of cytokine released. Spot size refers to relative amounts of cytokine produced per cell.

Although ELISpot assays are widely used in the context of the HIV study, the relationship between spot size or spot count and other variables has not been studied until now. For this reason we are interested in studying if there is a relationship between spot size or spot count, and other different clinical parameters, using different statistical techniques, such as balanced/unbalanced one-way/two way ANOVA (Sahai and Ojeda, 2004) or Generalized Linear Models (GLM). Moreover, as the immunospot reader creates an Excel sheet for every ELISpot assay, we have built a routine in \texttt{R} to read and sort the data and we are planning to create an \texttt{R} package for the statistical analysis of ELISpot assays data.

\textbf{Keywords:} ANOVA, ELISpot assays, Generalized Linear Models, HIV studies.

\textbf{References}

Is arsenic and cadmium exposure causally associated with renal damage? A Mendelian Randomization study

Maria Grau-Perez¹, Jose D. Bermudez Edo², Jose L. Gomez-Ariza³, Zoraida Soriano-Gil⁴, J.Antonio Casasnovas⁴, Josep Redon¹ and Maria Tellez-Plaza⁵

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Introduction: Traditional epidemiologic studies are limited to evaluate correlational associations due to confounding and other biases linked to observational data. Mendelian randomization (MR) is the use of genetic variation to assess the causal effect of modifiable exposures on health outcomes using observational data. Using a MR approach, we investigated the causal relationship of arsenic (As) and cadmium (Cd) exposure with renal damage among 1323 participants of the Aragon Workers Health Study (AWHS) from Spain.

Methods: Exposure to As and Cd was evaluated by its concentrations in urine. Renal damage was assessed as the urine albumin levels. We first conducted a genome-wide association analysis to identify single nucleotide polymorphisms (SNPs) associated with increased levels of urine metals in an independent database. Identified SNPs were further genotyped in the AWHS for this analysis. We validated 3 and 2 SNPs associated with genetically elevated urine As and Cd levels, respectively. We evaluated the correlational association of As and Cd urine levels with urinary albumin using traditional linear regression models and the causal association using a 2-stage least squares MR method.

Results: In traditional adjusted linear regression models, the geometric mean ratio (95% confidence interval) of urinary albumin by an interquartile range increase in As and Cd levels was 1.03 (0.97, 1.10) and 1.06 (1.01, 1.11), respectively. The F-statistic of the SNPs for As and Cd was 14.8 and 8.7, respectively, suggesting that the instrumental variables for Cd might be weak. The MR analysis suggested a significant causal association of As and Cd exposure with increased urinary albumin. The GMR (95% CI) of urine albumin per metal-increasing allele was 1.58 (1.15, 2.16) and 1.59 (1.18, 2.14) for As and Cd, respectively.
Conclusion: Mendelian randomization is an interesting tool to disentangle causal associations were randomized controlled trials are not feasible. More research is needed to confirm these findings.

References


estancia hospitalaria entre otros. El alcoholismo y el tabaquismo se presentan como factores de riesgo para la supervivencia a 5 años así como para el uso de servicios sanitarios y la calidad de vida de los pacientes en el mismo periodo.

**Conclusiones:** Los individuos con detección clínica y cáncer de intervalo presentan diferentes características respecto a los otros grupos, así como peores resultados clínicos, menores complicaciones y menor calidad de vida durante el tiempo de estudio. Además, la variabilidad entre los hospitales tampoco presenta grandes diferencias a lo largo del tiempo de estudio.

**Keywords:** cáncer colorrectal, modelos jerárquicos, servicios sanitarios.

**References**


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**A statistical model for analysing the effects of treatments on multivariate time series of microbiota data**

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There is an increasing interest on understanding the microbiota dynamics and predicting future microbiota behaviours which has motivated the development of dynamical system models. In particular, being able to predict with a model allows us to analyse the response of the microbiota to perturbations. The data available is the relative abundance of the families of bacteria of the microbiota of VIH patients. In this work, we analyse longitudinal microbiota data taking into account the compositional feature of the microbiota data. We assume that the data is distributed using the Dirichlet distribution and we present an autoregressive model that is able to predict the population dynamics. To make a long story short, we present a model that
enables quantifying the effects of treatments on microbiota time-series from VIH patients.

**Keywords:** time-series, autoregressive model, microbiota.

**References**


Big problems, integrated solutions

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In Ecology, due to the monitoring effort and the inaccessibility of breeding areas, it is often the case that a fraction of the population is monitored. These monitored areas are assumed to be a random sample of the whole population. However, this assumption is often violated. As a consequence, parameter estimates obtained from the monitored areas may be biased (Sanz et al., 2016). This problem is called partial monitoring.

We consider a particular case related to a large capture-recapture-recovery database obtained from the largest colony of Common guillemot (Uria aalge) of the Baltic Sea. We model annual survival and resighting probabilities in relation to the age of the individuals through integrated models (Besbeas et al., 2002). In particular, we establish four age categories: 1 (one year old individuals), 2 (two years old individuals), 3 (three years old individuals) and 4 (individuals from four to ten years old).

In this colony, the partial monitoring is present at breeding ledges, where mainly 3 and 4-10 individuals are resighted. As a result, survival probabilities for those age classes are underestimated while recapture probabilities are overestimated. In this study we compare the results obtained from the standard Cormack-Jolly-Seber model and the suitably parameterised integrated model that incorporates recoveries from all individuals in the colony (and not only those at the monitored sites). The results obtained show how the use of the integrated models solve this partial monitoring problem in this colony, although it can be easily extended to other populations.

References


Mixed models for exploring sardine fishing in the Mediterranean Sea

Gabriel Calvo¹, Rubén Amorós², Carmen Armero¹, Maria Grazia Pennino³ and Luigi Spezia⁴
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The European pilchard (Sardina pilchardus) is one of the most exploited small pelagic fish species. It plays an important role in the transfer of energy from lower to higher trophic level organisms but some worrying changes have recently been observed on its fishing. Experts assure that the abundance of this fish has decreased. In fact, in 2017 the International Council for the Exploration of the Sea advised to stop fishing them for the next fifteen years everywhere.

In this work, we explore the temporal evolution of sardine fishing in the Mediterranean Sea from 1970 to 2014 by means of a few Bayesian longitudinal models which include three types of random variation, i.e., random effects (in the intercept and slope), serial correlation (an autoregressive term) and measurement error. Model assessment is performed through exploration of the conditional and marginal residuals.

Keywords: Fishery estimation, longitudinal models, random variation, serial correlation.

From data-limited to data-rich: the evolution of sardine in the Bay of Biscay

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¹AZTI, Sustainable Fisheries Management
²BCAM, Applied Statistics

In the last years sardine (Sardina pilchardus) in the Bay of Biscay has evolved from being considered a data-limited stock to being classified by ICES (International Council for the Exploration of the Sea) in Category 2. This means the stock has analytical assessments and forecasts, but they are only treated qualitatively in relative terms. This raises additional difficulties, such as the calculation of reference points (i.e. maximum sustainable yield) and the definition of harvest control rules in relative terms. In this study we conducted management strategy evaluation for sardine in the Bay of Biscay to compare the performance of absolute and relative harvest control rules. All the analyses were done using FLBEIA, which is an R package for the development of bio-economic impact assessments of fisheries management strategies. The population dynamics were based on the latest ICES stock assessment and included uncertainty in the historical period and the biological parameters. Age structured and aggregated abundance indices from different surveys and age
structured observations from the catches were simulated and a stock assessment catch at age statistical model was explicitly incorporated into the simulation. The results will allow to initiate the discussion of the best management strategy for this stock.

**Keywords:** simulation study, population dynamics, uncertainty estimation, stock assessment models, management strategy evaluation.

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**Variable selection for genome wide association analysis in plant breeding**

Iker Oyanguren Monferrer$^1$ and Anabel Forte$^1$

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In modern plant breeding comparative experiments, there is often a necessity to accurately quantify the underlying genetic structure of industry driven phenotypic traits. With the availability of low cost high-throughput genotyping technology, this quantification usually involves the determination of quantitative trait loci (QTL) through a high dimensional genome wide association analysis pipeline. Within this pipeline, there may be several methods to reduce the dimension of the genetic marker set to contain the most important markers for further scrutiny. In this work we discuss a two steps genome wide association analysis which makes use of a Bayesian variable selection strategy to determine markers significantly linked to a supposed QTL. The method is illustrated with a large Australian wheat panel.

**Keywords:** QTL, Bayesian model selection, SUPER method.

**References**


Session 3: Survival analysis and Optimum experimental design
September 5th, 15:30 to 16:45
Chair: Yovaninha Alarcón Soto

Prognostic factors in the survival of women with breast cancer through a Bayesian competitive risk model

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The World Health Organization deals with breast cancer is the most frequent cancer among women, impacting 2.1 million women each year, and also causes the greatest number of cancer deaths among women. Breast cancer rates are higher among women in more developed regions, rates are increasing in nearly every region globally (WHO, 2018). The data of this study comes from a follow-up of 288 months to 1215 patients diagnosed with breast cancer in the region of Biobio in Chile, where 333 died of breast cancer, 121 due to another cause and 761 were still alive at the time of finalization of the study. The objective of this study is to identify those clinical variables (stage of the disease and age at the time of diagnosis, age at the time of menarche, and menopause, age at first birth, if there is a family history of cancer, etc), associated with the prognosis of the women survival with breast cancer. For this, Bayesian models of Cox’s and Cause-specific risks (Klein et al, 2014.,Schmorr et al, 2013) are used that consider the clinical variables of the study as covariates.

Keywords: Survival analysis; Measuring time scale; Censored observations.

References


Evaluación basada en la modificación de la historia natural de la efectividad de un programa poblacional de atención a la insuficiencia cardíaca

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5Equipo de Atención Primaria de Arrasate-Mondragón, Organización Sanitaria Alto Deba, Arrasate-Mondragón, Gipuzkoa.

El objetivo de este trabajo fue evaluar la efectividad en tres Organizaciones Sanitarias Integradas (OSIs) de un programa integrado de atención a la insuficiencia cardiaca (IC) desarrollado en el País Vasco.

La historia natural de los pacientes con IC se caracteriza por un proceso dinámico con periodos de compensación y descompensación en el que es habitual que los ingresos hospitalarios sean recurrentes y cada vez más frecuentes. Es por ello que las intervenciones dirigidas a la IC han sido comúnmente evaluadas mediante los ingresos hospitalarios de los pacientes.

Para la evaluación del programa se llevó a cabo un estudio observacional retrospектив basado en las bases de datos administrativas y clínicas de Osakidetza entre los años 2014 y 2018. Se midió el efecto de la intensidad de la intervención sobre el tiempo hasta ingreso por IC, ajustando por otras variables de interés. Para ello se utilizó el modelo de procesos de conteo de Andersen-Gill, una extensión del modelo de riesgos proporcionales de Cox para analizar no solo los primeros eventos de cada paciente, sino también las recurrencias. Estos modelos permiten la inclusión de variables tiempo dependientes en el análisis, lo cual nos posibilitó medir la intensidad de la intervención antes de cada hospitalización.

Se analizó un total de 6.768 pacientes. La edad (hazard ratio (HR)=1,016; IC95% 1,011-1,022), el índice de Charlson (HR=1,067; IC95% 1,047-1,087) y el número de hospitalizaciones previas (HR=1,632; IC95% 1,557-1,712) fueron factores de riesgo de reingreso. La intensidad de la intervención se asoció a una reducción de las hospitalizaciones. También se observaron diferencias entre OSIs, aunque el análisis de la interacción entre la OSI y la intensidad de la intervención mostró que estas diferencias desaparecen a medida que la intervención se intensifica.

Se concluyó que la efectividad de un programa educativo y de monitorización de atención a la IC depende de su nivel de implementación. Además, la evaluación de las intervenciones para la IC basada en los reingresos debe tener en cuenta toda la historia natural de la IC, para lo cual resultan de utilidad las técnicas de análisis de la supervivencia para eventos repetidos.

Keywords: insuficiencia cardiaca; programa integrado; análisis de la supervivencia; eventos repetidos; modelos Andersen-Gill.
Statistical Modeling in sports injury prevention

Lore Zumeta-Olaskoaga\(^1\) and Dae-Jin Lee\(^1\)

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Sports biostatistics is an emergent field in the Sports Science area. Initially proposed by Casals and Finch (2018), the role of the sports biostatistician brings together statistics, epidemiology and sports medicine. Sports science is not exempt from statistical applications and here one of the principal interests relies on the prevention of sports injuries. Statistics can help in gaining insights and understanding why sports injuries occur.

In this talk, a number of important issues regarding the complex nature of sports injury data will be presented. Some statistical aspects when quantifying the injury risk such as measures of injury epidemiology will be shown. Survival analysis will be introduced as a tool to appropriately handle this type of data, identify the main risk factors and model injury occurrence. Also, real-world examples will be used to illustrate these concepts. We will end up looking over the methodological challenges that sports biostatistics currently faces.

Keywords: sports injury, sports biostatistics, survival analysis.

References


Pensar antes de actuar: introducción al diseño óptimo de experimentos

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En la actualidad existe un interés creciente en el estudio del diseño de experimentos, ya que es una parte fundamental del método científico. El análisis de los datos será informativo sólo si los datos por sí mismos lo son (1). Las ciencias experimentales y la ingeniería son campos del conocimiento especialmente interesados en obtener modelos que expliquen adecuadamente los fenómenos objeto de estudio. La obtención de estimadores precisos de los parámetros del modelo es, entre otras, una propiedad deseable para obtener la mejor calidad de la inferencia estadística. Por este motivo, la estrategia de recogida de datos se convierte en un punto crucial para el buen desarrollo del estudio donde factores económicos y limitaciones prácticas entran en juego. El principal objetivo del Diseño Óptimo de Experimentos (DOE) consiste en determinar dónde tomar las observaciones y con qué frecuencia para optimizar algún aspecto del modelo de manera eficiente. En este trabajo se presenta una introducción general a esta teoría, poniendo en valor algunos de los modelos abordados por el grupo Optimum Experimental Design (2) desde esta perspectiva. Algunos de ellos son utilizados para la calibración de instrumental utilizado en radioterapia, para explicar el efecto de una droga en la muerte de células tumorales, para detectar el fenómeno de hormesis o para análisis de supervivencia. Finalmente, se proporcionan diferentes ejemplos prácticos en los que se comprueba cómo un estudio y una planificación previa mejora notablemente la precisión de la inferencia estadística, aprovechando la máxima información recogida de cada unidad experimental realizada.

**Keywords:** Diseño óptimo de experimentos, inferencia estadística

**References**


Three-level hierarchical modelling of longitudinal data: Application to the study of worldwide obesity prevalence

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This work considers hierarchical or multilevel curve modelling for longitudinal data. Motivated by the need for analyzing the evolution of the difference in the prevalence of obesity between males and females, we propose a three-level nested hierarchical model. In particular, our interest focuses on modelling and estimating the evolution of the difference in the prevalence of obesity from 1975 to 2016, and we have information for 187 countries nested into 10 regions, and regions into the world population (Gender Statistics, The World Bank, https://datacatalog.worldbank.org/dataset/gender-statistics). Since important decisions may need to be taken at each level of the hierarchy (countries, regions or worldwide), estimation must be as accurate as possible. Thus, a model which can handle appropriately with hierarchically nested effects is needed. Hierarchical semiparametric and nonparametric regression methods can be found in the literature. Examples include the use of smoothing splines, penalized splines or functional principal component (e.g., Durbán et al., 2005; Fitzmaurice et al., 2011; Brumback and Rice, 1998; Li et al., 2015). In this work, we propose an extension of the spline-based two-level nested hierarchical approach described in Brumback and Rice (1998) and Durbán et al. (2005), among others. We propose the use of penalized splines (P-splines) jointly with second-order differences (Eilers and Marx, 1996) and its representation as a mixed-effect model (Currie and Durban, 2002). This approach implies that each function in the hierarchy is composed by a first order polynomial (modelled as fixed or random, depending on the level) plus a nonlinear effect. Estimation is based on the SOP method presented in Rodríguez-Álvarez et al. (2019). We compare our results with the ones obtained using the methods implemented in the R-package mgcv (Wood, 2011) and using regression with functional data through the R-package refund (Goldsmith et al., 2018).

Keywords: P-splines, hierarchical model, longitudinal data, prevalence of obesity.

References


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**INLAMSM R-package: Multivariate spatial models using R-INLA.**

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Literature about multivariate spatial models have increased in the last years due to the fact that classical univariate methods are not able to consider the real cohabitation of different individuals or procedures (for instance, different diseases in a disease mapping context). Bearing in mind that multivariate procedures lead to high computation time procedures, different ideas have been developed to speed up fitting multivariate spatial models. On the one hand, some authors have re-parametrized the classical generalizations of the univariate models in order to avoid high computational-time operations within the framework of MCMC algorithms (Botella-Rocamora et. al. 2015). On the other hand, some authors prefer approximate methods, such as the INLA method (Rue et. al. 2009) which is commonly used in order to avoid the high computation time of the MCMC models. This idea is part of the motivation of this work whose main goal is to implement, using R-INLA benefits, multivariate areal models which were proposed for been computed using MCMC algorithms and then re-parametrized in order to be more computationally efficient. Moreover,
comparisons with other existent methodologies have been carried and all the methods implemented have been gathered in an R-package named INLA Multivariate Spatial Models: INLAMSM.

**Keywords:** INLAMSM, Bayesian statistics, R-INLA, Spatial models.

**References**


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**CompARE: a web app to study Composite Endpoints**

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**Context:** Composite endpoints are frequently chosen as primary endpoint in randomized controlled trials. Composite endpoints are common in many areas, such as cardiovascular and oncology, where Major Adverse Cardiovascular Events (MACE) and Progression Free Survival (PFS) are the usual primary endpoints. The choice of the primary endpoint, together with the corresponding needed sample size, is critical for the trial success, but deciding which components should be included in the composite endpoint is not straightforward. We have developed an Asymptotic Relative Efficiency (ARE) measure as an aid for this decision. ARE depends on the marginal and joint distributions of the composite components.

**Objective:** To provide a tool for researchers in studies including several time-to-event or binary outcomes. CompARE, in a quantifiable and friendly way, allows to select the components of a primary composite endpoint and its corresponding sample size and to study the attributes of the composite treatment effect.

**Methods:** We present the web-app CompARE (https://cinna.upc.edu/compare). The user can specify the probabilities of event-occurrence; the expected treatment effect through hazard ratios or odds ratios; and the degree of association between components. As a result, CompARE provides: 1) the ARE values depending on different treatment effects and degrees of association by means of tables and plots; 2) sample size calculations under several scenarios; and 3) some treatment effect measures for the composite endpoint.

**Results:** We present CompARE for the ZODIAC and TACTICS-TIMI 18 trials with PFS and MACE as primary endpoint, respectively. Through these examples, we illustrate how to decide whether is more efficient to use the composite endpoint.
or to remain with one of the components; to compute the sample size under different scenarios and to learn about the impact of the association between components on the composite endpoint. We use CompARE to discuss about the proportionality hazards assumption of the composite endpoint in the ZODIAC trial or the different behavior of relative and absolute effect measures in the TACTICS-TIMI 18 trial.

**Conclusions**: CompARE is a valuable and helpful tool for the discussion of the primary endpoint to lead the randomized controlled trials regardless whether you have binary or time-to-event components.

**Keywords**: Software, Clinical Trials, Survival Analysis.

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**References**


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**Fitting Bayesian Dirichlet regression using the integrated nested Laplace approximation**

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Dirichlet regression models can be used to analyze a set of variables lying in a bounded interval that sum up to one exhibiting skewness and heteroscedasticity, without having to transform the data. These data which mainly consist of proportions or percentages of disjoint categories are widely known as compositional data and are common in areas such as ecology, geology, and psychology. Bayesian inference has become a popular tool to deal with complex models such as Dirichlet regression models. The integrated nested Laplace approximation (INLA) is a widely extended methodology that for a large class of models provides higher accuracy for a limited computational budget. However, the implemented R-INLA package can not deal with multivariate likelihoods, such as, in particular, the Dirichlet likelihood. In this work, we propose an expansion of the INLA method for Dirichlet regression.

**Keywords**: Dirichlet regression, INLA, multivariate likelihood.

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**References**

Regression models for count data often present overdispersion, a phenomenon that arises when the real variance of the data is larger than the one specified in the model. In addition, when working with georeferenced data, the spatial dependence that may exist among the different locations must be also taken into account in order to produce reliable inference processes from the estimations. In this work we review spatial conditional overdispersion models associated to the binomial distribution, such as the spatial conditional beta binomial and binomial normal models. In these models, spatial association structures are specified in the regression model for the conditional mean. This specification allows researchers to be able to quantify the intensity of the spatial association that may be present in the data. Besides, assuming that the overdispersion is partially generated by the spatial autocorrelation in the data, the model would also be able to capture this phenomenon. We analyze the number of mother who had given birth to their last child between 1999 and 2005 and who underwent a postnatal screening period in different departments in Colombia, and study the behavior of this variable as a function of some specific covariates available in this dataset, illustrating the usefulness of the proposed models. Additionally, we also assess the performance of such models when including a conditionally autoregressive (CAR) spatial random effect in the model specification. Models have been fitted with the use of the Markov Chain Monte Carlo (MCMC) algorithms within the context of Bayesian estimation methods.

**Keywords:** Overdispersion models, Spatial models, Bayesian methods.

**References**


Design of an alternative delimiting strategy for the *Xylella fastidiosa* infection in Alicante: a development of a sequential adaptive algorithm for optimising survey and sampling.

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*Xylella fastidiosa* (*Xf*) is a regulated quarantine plant pathogen in the European Union (EU). The current legal provisions specify the implementation of delimiting surveys in infested areas to demarcate the geographic extent and implement eradication or containment measures (EU, 2015). North-eastern Alicante Province, Spain, is one of the areas infested by *Xf* in the EU, and so it is currently demarcated and subjected to a delimiting survey activity. Based on the 2018 official surveillance program, in which approximately 100,000 ha. were surveyed and 11,000 samples were taken and analysed, this work aimed to improve the efficiency of the current delimiting survey strategy.

A generic delimiting strategy was designed based on a three-phase adaptive approach in which the surveyed area (epidemiological unit) size and sampling intensity were tailored according to the previous phase information (Peyrard et al., 2013). An algorithm was implemented to optimise the number of epidemiological units to be surveyed and the sample size by simulating different random sampling scenarios from the reference data. Strategy evaluation was performed by comparing the delimitation efficacy and incidence estimates between the proposed and the current strategy. Incidence was modelled by means of a Bayesian spatial hierarchical model in which climatic and spatial factors were evaluated using INLA approximation (Lindgren and Rue, 2015). The proposed delimiting strategies achieved to delimit similar extension disease (similar efficacy) with a lower number of samples (better efficiency) than current.

**Keywords:** spatial sampling, adaptative sampling, simulations-optimization.

**References**

Assessing landscape fire hazard by multitemporal automatic classification of Landsat time series using Google Earth Engine in West-Central Spain

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Annual Land Use and Land Cover (LULC) maps are needed to identify the interaction between landscape changes and wildland fires. Objectives: In this work, we determined fire hazard changes in a representative Mediterranean landscape, through the classification of annual LULC types and fire perimeters, using a dense Landsat Time Series (LTS) during the 1984-2017 period, and MODIS images. Methods: We implemented a semi-automatic process in the Google Earth Engine (GEE) platform to generate annual imagery free of clouds, cloud shadows, and gaps. We compared LandTrendr (LT) and FormaTrend (FT) algorithms that are widely used in LTS analysis, to extract the pixel tendencies and, consequently, assess LULC changes and disturbances such as forest fires. These algorithms allowed us to generate the following change metrics: type, magnitude, direction and duration of change, as well as the pre-change spectral values. Results and conclusions: Our results showed that the FT algorithm was better than the LT algorithm at detecting low-severity changes caused by fires. Likewise, the use of the change metrics’ type, magnitude, and direction of change increased the accuracy of the LULC maps by 4% relative to the ones obtained using only spectral and topographic variables. The most significant hazardous LULC change processes observed were: deforestation and degradation (mainly by fires), encroachment (i.e., invasion by shrublands) due to agriculture abandonment and forest fires, and hazardous densification (from open forests and agroforestry areas). Although the total burned area has decreased significantly since 1985, the landscape fire hazard has increased since the second half of the twentieth century. Therefore, it is necessary to implement fire management plans focused on the sustainable use of shrublands and conifer forests; this is so because the stability in these hazardous vegetation types is translated into increasing fuel loads, and thus an elevated landscape fire hazard.

Keywords: FormaTrend; Landsat imagery; land cover changes; LandTrendr; landscape fire hazard; time series analysis; random forest.
La validación de los nuevos inmunoensayos que se desarrollan consiste en evaluar si ciertas propiedades de éstos se cumplen. Una de ellas, y que sólo aplica a los métodos cuantitativos, es la de la linealidad de la exactitud del método. La linealidad se refiere a la capacidad que tiene el método de reportar resultados que son directamente proporcionales a la concentración de analito en la muestra, es decir, $y = kx$.

Esto hace pensar que el modelo de regresión lineal estimado mediante el método de mínimos cuadrados sea una buena estimación de los datos. Sin embargo, debido a que los datos procedentes de inmunoensayos, por lo general, presentan heterocedasticidad y autocorrelación, se ha decidido analizar si determinados métodos mejoran la estimación lineal (en el sentido de que la estimación presente menor varianza) con respecto al de los mínimos cuadrados. Los métodos propuestos han sido: (1) regresión ponderada, (2) transformación estabilizadora de la varianza y (3) los estimadores de Theil - Sen y Siegel.

La evaluación de las diferentes técnicas estadísticas se ha llevado a cabo mediante la simulación de datos según 4 patrones de la varianza ampliamente observados en datos procedentes de inmunoensayos. Estos patrones son: (1) varianza constante, (2) varianza lineal, (3) coeficiente de variación constante y (4) varianza con forma de gancho. La comparación de estos métodos estadísticos también ha sido evaluada sobre datos reales para verificar que se obtenían resultados similares.

Finalmente, se ha visto que de todos los métodos estudiados el estimador de Theil - Sen es el que presenta menor varianza, sin embargo no hay una gran diferencia con la estimación obtenida mediante el método de mínimos cuadrados, por lo que con esta estimación no se puede garantizar la existencia de mejores resultados en la evaluación de la linealidad.

Igualmente, se ha elaborado un protocolo que sirve de guía a los investigadores que han de realizar estos estudios de validación.

**Keywords:** Heterocedasticidad, regresión lineal, inmunoensayos, rango de linealidad, curva maestra.

### References


Benefits and drawbacks of using stratification factors in clinical trials. When and why should we use it?

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The inclusion of stratification factors is increasingly common in clinical trial design, especially in phase III studies. This is explained by two main reasons: i) to prevent imbalance in randomization across known prognostic factors; and ii) to provide a more accurate estimation of treatment effect.

Several techniques exist to control the randomization process according the stratification factors: block randomization and dynamic allocation are the most commonly used. However, some concerns arise when the number of strata increase and, consequently, the risk of imbalance and the probability of empty strata become a real problem. In time-to-event analysis the stratified Cox model (an extension of Cox model) is used to estimate the treatment effect. On one hand, this methodology provides a more accurate point estimation if stratification factors are, in fact, prognostic factors. On the other hand, the estimate standard deviation becomes bigger as the number of strata increase. Under this scenario, several open questions arise in the context of study design. What is the cost of over-stratification? How many prognostic factors to be included as stratification variables? How predictive factors should be treated in study design?

The goal of the current study is to analyse all these questions via simulation under different scenarios. Evaluation of type I/II errors will be studied in detail for each scenario to estimate their consequences for the final treatment effect estimation.

Keywords: Stratification factor, Clinical trial, Randomization, Over-stratification

Robustness of design for accelerated failure time models with Type I censoring

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Accelerated Failure Time (AFT) models are fairly commonly used in the field of manufacturing, but they are more and more frequent for modeling clinical trial data. These models are defined through the survival function of the time-to-event variable, T. This work deals with construction of robust designs for AFT models with the possibility that the Acceleration Factor (AF) is misspecified when the variance of T is known. In particular, we allow the “true” AF to vary over a neighbourhood of possible functions, \( AF(x, \theta) = \exp(\theta' x + g_n(x)) \), for some unknown perturbation function \( g_n \). As in Huber (1975), we consider \( g_n(x) \) lies in a contaminating space \( F \). Thus, we cannot asses the efficiency of a design through the covariance matrix of the Maximum Likelihood Estimator (MLE). In this case, the estimate is subject
to both “bias error” due to the inadequacy of the model, as well as “variance error” due to sampling. We obtained the asymptotic Mean Squared Error matrix (MSE) of parameter estimates for right censored observations. In order to the asymptotic treatment we will assume that $g_n$ functions are of the order $o(1/\sqrt{n})$. Finally, a typical distribution in AFT, Log-Logistic distribution, was considered to apply the above results.

**Keywords:** Optimal Experimental Design, Survival Analysis, Robustness, Accelerated Failure Time Models.

**References**


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**Efficiency Analysis for the Misspecification Problem of Response Probability Distribution in Optimal Experimental Design**

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It is a common assumption in the context of Optimal Experimental Design that the response variable follows a homoscedastic normal distribution. There are, however, other studies that assume different probability distributions based on prior experience or additional information. The main goal of this study is to look at the effect, in terms of efficiency, of misspecification in the probability distribution on optimal design. From the elemental information matrix, which includes information on the probability distribution of the response variable, a generalized Fisher information matrix is obtained. Relevant theoretical results were obtained, for different regression models, comparing heteroscedastic Poisson, gamma and normal distributions. Finally, the analysis was broadened to include a practical case which considers a 4-parameter Hill model, to explain the effect of a pharmaceutical drug on cell development.

**Keywords:** Elemental Information Matrix; Gamma Distribution; Poisson Distribution; D-optimization; D-efficiency.
In an increasingly globalized world, it seems obvious that digital technology has transformed society. Modern society is immersed in a continuous process of knowledge accumulation and new developments. In turn, economic interdependence is growing between areas and less developed regions are being left behind by structural changes in the world economy. This phenomenon has positive effects such as greater development either economic growth, nevertheless at the same time it requires a certain technical level incorporated into the productive tasks and a high-performing human capital applying this new knowledge. Likewise, the development of innovations and improvements in productive activity are part of the market competitive strategy.

The European Union, aware of international influence, considers innovation as the main course of action for smart and sustainable growth. Technological advances have revealed the transcendence of knowledge as the dominant intangible asset. In fact, knowledge creates Intellectual Capital, an asset that can provide a competitive edge. Intellectual Capital, therefore, should be considered an economic investment, just as capital formation or labour.

Normally, productive performance is analysed on a national level; however, what is significant for the whole may not be for parties, and that is why a regional analysis is highly valuable for the diagnosis of economic functioning. The goal is to present a proposal for intellectual valuation at regional level and to know how its effect on productivity can be channelled. The analysis is based on the experience of 271 regions NUTS 2 in Europe by 2015. Through the perspective of intellectual capital, useful elements are provided to observe and understand the behaviour of productive structures in Europe in the face of the digitalisation process and discover singularities and inequalities between regions.

Keywords: human capital, technology, innovation, investigation, productivity.
Aplicación de la Teoría de Respuesta al ítem para armonizar diferentes escalas de depresión en el proyecto ATHLOS

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La armonización de datos es una técnica que sirve para lograr o mejorar la comparabilidad de variables y medidas similares recopiladas en estudios diferentes. Armonizar variables como sexo, edad, altura, entre otras, es relativamente sencillo ya que consiste en recodificaciones de categorías o cambio de unidades. En cambio, para variables medidas a través de escalas psicométricas, como por ejemplo depresión, este proceso resulta más complejo dado que para un mismo constructo pueden existir varias escalas con características psicométricas distintas. Por ello, armonizar la variable directamente de la escala propia de cada estudio podría llevar a un sesgo de medida entre los estudios. Un proceso alternativo sería la construcción de una nueva escala con síntomas depresivos armonizados entre los diferentes estudios y la aplicación de modelos de variables latentes.

En la presentación se mostrará la aplicación de este proceso alternativo de armonización usando los datos de las escalas de depresión de 14 estudios longitudinales que se armonizaron en proyecto ATHLOS (Ageing Trajectories of Health: Longitudinal Opportunities and Synergies) con un tamaño de la muestra final de más de 200.000 individuos. En este proceso, primero se seleccionaron 12 ítems que fueron considerados los más comunes e importantes, extraídos de las siguientes escalas de depresión: EURO-D, CIDI, BDI-II, GHQ12 y CES-D versión de 8, 10 y 20 ítems. A continuación se estudió la unidimensionalidad del constructo de depresión utilizando el análisis factorial exploratorio y confirmatorio. Seguidamente, ajustando un modelo de la Teoría de Respuesta al ítem, se calcularon las puntuaciones para cada individuo de manera continua. Por último, se usó la curva ROC para encontrar el punto de corte óptimo para su dicotomización entre individuos con depresión y sin depresión.

Keywords: Armonización, Teoría de Respuesta al ítem, depresión, escalas psicométricas

References


Caracterización del dimorfismo sexual a nivel craneal mediante análisis discriminantes

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El dimorfismo sexual es el principal patrón de variación biológica entre individuos adultos dentro de las poblaciones humanas actuales. El componente esquelético juega un papel clave en la determinación del sexo en campos como la Antropología o las Ciencias Forenses. No obstante, dentro del conjunto del esqueleto humano, no todos los huesos presentan el mismo grado de dimorfismo, y el cráneo es considerado por detrás de la pelvis, como la estructura más dimórfica. Estudios previos han caracterizado ya algunas de las estructuras craneales que presentan mayor dimorfismo, entre ellas se encuentran los arcos superciliares, las apófisis mastoides o el hueso zigomático. El presente estudio tiene como objetivo caracterizar el dimorfismo sexual a nivel craneométrico. Los datos analizados corresponden a la base de datos de Howells que comprende un total de 2524 cráneos (1368 varones y 1156 mujeres) pertenecientes a 30 poblaciones mundiales diferentes, y de los cuales se tiene información de un total de 46 variables craneométricas cuantitativas pertenecientes tanto al neurocráneo como al esplacnocráneo y consistiendo tanto en medidas de anchura, como de altura o longitud. El análisis de los datos se llevó a cabo mediante el software estadístico R. La metodología empleada para determinar qué variables craneométricas explican mejor este dimorfismo sexual comprendió un Análisis de Componentes Principales (ACP) seguido de un Análisis Discriminante (AD) para optimizar su caracterización. Entre los principales resultados obtenidos el índice craneal, el cual refiere la forma general del cráneo, no mostró un peso relevante a la hora de caracterizar el dimorfismo sexual, siendo éste por tanto más debido al tamaño y no a la forma del cráneo. Además, el ACP indicó que las variables de anchura facial son las que mejor explican la separación de los individuos por sexos. Este fenómeno se puede contrastar y explicar mediante la bibliografía, relacionando una mayor anchura facial en los hombres con posibles efectos de selección sexual. El AD indicó que la variable más dimórfica es la anchura bicigomática. La precisión alcanzada por el modelo a la hora de clasificar los individuos correctamente por sexos fue del 92,24%. Este estudio muestra la utilidad de las variables craneométricas cuantitativas para discriminar el sexo del individuo mediante la construcción de funciones discriminantes.

Keywords: Craneometría, Dimorfismo sexual, Análisis Discriminante.
From high-dimensional to functional data: application in SNPs data

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Stringing via manifold learning is introduced, a method that allows to treat high-dimensional data as functional data. It takes advantage of the high dimensionality of the data by considering them as discretized observations of some underlying smooth stochastic process. Assuming that data is observed with a scrambled order of its components, stringing starts with a reordering step and then maps the high-dimensional vectors into functions. We extend the earlier version of stringing (Chen, Müller, and Wang, 2011), based on multidimensional scaling, by incorporating manifold learning. Our method is able to recover non-linear relationships between predictors, resulting in a better reordering of the data and a more reliable functional representation. This work reviews the stringing methodology and discusses the techniques related to it. Through simulation studies it is shown that our contribution outperforms the previous approach. We remark the usage of functional principal component analysis through conditional expectation, as a useful tool for functional representation when data is rather noisy. Illustrative examples regarding stringing and functional data representation are provided.

A real data application on graft-versus-host-disease (GVHD) (Martínez-Laperche et al., 2018) is also addressed. The study deals with patients who had an allogeneic hematopoietic stem-cell transplantation (allo-SCT) in order to treat hematologic malignancies. GVHD is considered the main complication after allo-SCT (affects 30% – 50% of patients) and there are not validated laboratory tests to predict it. Stringing via manifold learning is applied to single-nucleotide polymorphisms (SNPs) data, providing functional genotypes of ill and healthy patients.

**Keywords:** Functional data analysis, High-dimensional data, Functional principal components analysis, Multidimensional scaling, Manifold learning, Isometric feature mapping, Locally linear embedding; Functional genotype, SNPs data.

**References**


Overview and new proposals in the development of prediction models with complex survey data

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Nowadays, complex survey data are gaining importance between researchers in different fields. The sampling weights are one of the most remarkable differences between simple random sampling data and data collected based on a complex sampling design. Each sampling weight is the number of individuals that one observation represents in the population. However, despite the growing importance of these type of data, we believe that there is still a lack of statistical methodology to develop in this field. The first goal of this work is to make a review of the literature in the context of the development of prediction models with complex survey data.

One of the most discussed topics in this context has been the importance of incorporating the sampling weights in order to obtain accurate estimations of the parameters of prediction models. Specifically, in the context of logistic regression models for dichotomous response variables, the pseudo-likelihood function has been proposed as a modified version of the likelihood function that incorporates the sampling weights. But in addition to the estimation of the model parameters, there are many aspects that should be taken into account in the development of prediction models in order to end up with a good validated model: the selection of the predictors, the functional relation between outcome and predictors, imputation of missing values and evaluation of calibration and discrimination, among others. We believe that the sampling weights should be taken into account in throughout this process, beyond the estimation of the model parameters. In particular, the goals of this work are two fold: a) we focus on the impact of the sampling weights in the imputation process of a relevant predictor variable and, b) we propose a new methodology to impute the missing values of a dichotomous covariate in the context of complex survey data.

Keywords: complex survey data, prediction models, sampling weights.

References


Estudio comparativo de métodos de estimación de los parámetros del modelo de regresión logística con datos de encuestas con diseño complejo

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El modelo de regresión logística, es uno de los modelos predictivos utilizados con mayor frecuencia a la hora de analizar el efecto de diversas variables explicativas en una variable dicotómica. Cuando los datos provienen de una muestra aleatoria simple, los parámetros del modelo se estiman generalmente mediante el método de máxima verosimilitud. Sin embargo, cuando se dispone de datos provenientes de una encuesta basada en diseño complejo, cada individuo de la muestra tiene asociado su correspondiente peso muestral, que representa el número de individuos representados en la población. El propósito de este trabajo ha sido analizar el efecto que los pesos muestrales pueden tener -y el alcance del mismo- en la estimación de los parámetros del modelo de regresión logística. Para ello, se han empleado datos simulados basados en los datos de dos encuestas realizadas por el Instituto Vasco de Estadística (Eustat): ESIE (Encuesta sobre la Sociedad de la Información en las Empresas) y PRA (Encuesta de población en relación con la actividad). En ambas encuestas se ha realizado un muestreo estratificado de una etapa, estando los estratos definidos en base a una combinación de 3 variables en el primer caso, y una única variable en el segundo. El estudio de simulación consiste en la obtención de 200 muestras obtenidas a partir de un muestreo estratificado de una etapa. En cada una de estas muestras, se han estimado los parámetros del modelo de regresión logística utilizando dos alternativas: a) máxima verosimilitud, y b) máxima verosimilitud ajustada por los pesos muestrales. Los parámetros obtenidos se han comparado con los parámetros teóricos calculados en la población. Los resultados obtenidos muestran que las estimaciones obtenidas teniendo en cuenta los pesos de muestreo, se aproximan más a los parámetros teóricos. A pesar de que esto ocurre en ambos casos (ESIE y PRA), en el caso de ESIE las diferencias entre los dos métodos de estimación son mayores. Esto se debe a que el diseño del muestreo, y por tanto los estratos considerados en ESIE, son más complejos. Por lo tanto, los resultados del estudio de simulación sugieren que los pesos de muestreo han de ser considerados en el proceso de estimación de un modelo de regresión basado en datos de encuestas con diseño complejo.

Keywords: muestreo complejo, regresión logística, pesos de muestreo.

References


Targeted lipidomic profiling reveals a lipid mediator signature specific of acute-on-chronic liver failure

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Individuals with liver failure, severe disruption of the intrahepatic arterial and venous flow, diffuse fibrosis, and portal hypertension are diagnosed with cirrhosis. Traditionally, cirrhosis has been classified as compensated and decompensated.

In patients with cirrhosis or underlying chronic liver disease, Acute-on-chronic liver failure (ACLF) defines a syndrome characterized by acute and severe hepatic abnormalities. However, in contrast to decompensated cirrhosis, ACLF has a high short-term mortality. In 40-50% of the cases it is not possible to identify the cause of ACLF and in the remaining patients, active alcoholism and relapse of chronic viral hepatitis are the most common precipitating factors.

Lipid profiles from the CANONIC study, a prospective observational study including patients with acute decompensated cirrhosis (AD) and acute chronic liver failure (ACLF) were analyzed to assess the added value in risk and prognosis of lipid profiles to the available clinical data.

Non-parametric tests, Kruskal-Wallis test and Wilcoxon test were used to assess the association of the lipids with the disease status and organ failures. Age and gender were regressed out. Two types of logistic regression models to predict 6-month mortality were computed: including only clinical variables and including both lipid profiles and clinical variables were. Data was divided in two non-overlapping sets: training set including 80% of the samples and a validation set with the rest. Statistically Equivalent Signatures (SES) algorithm was used to select the “minimal” sets of features with the highest predictive power.

From 59 lipids analyzed, 15 show significant differences between healthy and AD or ACLF patients. Only two lipids show significant differences between AD and
A CLF. This tendency was seen also at follow-up. No significant added value of lipids data to the predictive models was observed.

**Keywords:** cirrhosis, acute decompensated cirrhosis (AD), Acute-on-chronic liver failure (ACLF), logistic regression, feature selection algorithms.

**References**


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Early-life urban exposome and cognitive and psychomotor function in children in 5 European birth cohorts: applying an exposome-wide association study approach

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**Introduction.** The exposome is a new paradigm that encompasses the totality of human lifelong environmental exposures, including the environmental urban exposures. Several regression-based statistical methods have been developed to assess exposome-health associations. One of the most used is the exposome-wide association study (ExWAS) approach, an exposure-by-exposure method accounting for multiple hypotheses testing.

**Objective.** We investigated the association between early-life urban exposome and cognitive and psychomotor function in children, applying an ExWAS approach.
**Methods.** We used data of 11,693 mother-child pairs from 5 European birth cohorts in Spain, France, Greece, UK, and Norway. Information on 30 exposures including air pollution, noise, climate, natural space, and built environment were estimated at each participant’s home address during pregnancy. Verbal, non-verbal, and psychomotor function were assessed using validated tests at 3-5 years old. We applied an ExWAS approach to study the association between the exposome and each outcome. We fitted linear regression models independently for each exposure and each outcome. The statistical significance of each association was assessed on the two-sided p-value after correction for multiple testing. Exposures found statistical significant were included in a multiple linear regression model and were retained if their two-sided p-value was below 5%. Models were adjusted for various socio-economic and lifestyle characteristics.

**Results.** Land use mixture was related to lower verbal function (-0.018 points (95%CI -0.036, -0.001)), while higher population density was related to higher verbal function (0.017 points (95%CI 0.001, 0.033)). Higher humidity levels were associated with lower non-verbal function (-0.081 (95%CI -0.153, -0.044)). Higher distance to green spaces was associated with lower gross motor function (-0.028 (95%CI -0.056, -0.001)) while higher particulate matter exposure was related to higher gross motor function (0.040 (95%CI 0.001, 0.079)). Higher particulate matter exposure was associated with lower fine motor function (-0.045 points (95%CI -0.078, -0.012) and -0.048 points (95%CI -0.093, -0.002), respectively). None of the associations survived correction for multiple testing, thus no multiple linear regression models were performed.

**Conclusion.** We found that air pollution, natural space, and built environment exposures during pregnancy were associated with small changes in cognitive and psychomotor function in children. We used an ExWAS approach, one of the most used methods developed to assess exposome-health associations. However, alternative methods such as Deletion/Substitution/Addition algorithm, an iterative selection method that selects the most predictive exposures of the outcome by cross-validation, taking into account

**Keywords:** ExWAS, multiple testing, exposome.
MESA REDONDA: “EL PAPEL DE LAS MUJERES EN LA ESTADÍSTICA”
September 6th, 12:00 to 13:30


Irantzu Barrio: Universidad del País Vasco (UPV/EHU). Irantzu Barrio es profesora adjunta del Departamento de Matemáticas Aplicadas y Estadística e Investigación Operativa de la Universidad del País Vasco UPV/EHU. Se licenció en Matemáticas en 2006 y obtuvo el Máster en Modelización Matemática, Estadística y Computación en 2010 en la misma universidad. Tras varios años trabajando como estadística en la empresa privada, en el año 2012 empezó a trabajar como docente en el Departamento de Matemática Aplicada, Estadística e Investigación Operativa de la UPV/EHU. En 2015 se doctoró en Matemáticas en la UPV/EHU. Su principal interés investigador se centra en el desarrollo de metodología estadística orientada a la resolución de complejidades derivadas de la investigación científica, principalmente en áreas biomédicas o experimentales. Los temas específicos de interés son el desarrollo y la validación de modelos predictivos. Además, trabaja en el soporte metodológico a investigadores tanto dentro de la facultad de ciencia y tecnología de la UPV/EHU como de unidades de investigación de centros del entorno (centros de investigación y hospitales) y el Instituto Vasco de Estadística (EUSTAT).

Anabel Forte: Universitat de Valencia (UV). Anabel Forte Deltell es profesora Contratada Doctora en el departamento de Estadística e Investigación Operativa de la Universitat de València. Entre sus líneas de investigación destaca la Selección de Modelos Bayesianos, así como el estudio de modelos estadísticos para datos correlados. En estas áreas ha publicado un total de 20 artículos Peer Reviewed además de varios artículos en proceedings y un capítulo de libro gracias a su participación en 8 proyectos competitivos financiados tanto por el gobierno de España como por el de la Generalitat Valenciana. En el ámbito de la docencia y la transferencia del conocimiento, Anabel ha participado en diversos proyectos de innovación docente y de divulgación. Ha impartido cursos en distintas universidades y realizado estancias de docencia en la Universitat Jaume I de Castellón y en la Universidad do Miño en Gimarães (Portugal), esta última financiada con fondos del programa Erasmus+. Ha dirigido dos tesis doctorales además de diversos trabajos fin de grado y máster dentro del grado en Matemáticas y del máster en Bioestadística de la Universitat de València. Además, es vocal de la junta directiva de la Sociedad Española
María Durban: Universidad Carlos III de Madrid (UC3M). María Durban es Almeriense, estudió su Licenciatura en Matemáticas en la Universidad de Granada (1993) y decidió emprender sus estudios de postgrado en el Reino Unido donde realizó un Máster en Estadística Matemática en la Universidad de Cambridge (1995), allí es donde experimentó por primera vez lo que realmente era la Estadística Aplicada y esto marcó su carrera desde entonces. A continuación, realizó sus estudios de doctorado en la Heriot-Watt University (Edimburgo, 1999) trabajando en la estimación de tendencias espaciales en enfermedades de plantas. Durante su experiencia postdoctoral en Biomathematics and Statistics Scotland (Edimburgo, 2000) trabajó en temas relacionados con las series temporales multivariantes aplicadas a la predicción de lluvia. Los años de formación en UK fueron para ella determinantes, no solo por la formación recibida, sino por la posibilidad de conocer de cerca a importantes Estadísticos/os que marcaron su carrera y con los que a día de hoy continúa colaborando. En el año 2000 se incorpora como profesora visitante a la Universidad Carlos III de Madrid donde en la actualidad es Catedrática de Universidad. Casi desde el inicio, su área de investigación se ha centrado en los modelos de regresión no-paramétricos, y en particular en los P-splines, en su formulación como modelos mixtos y sobre todo en su aplicación al análisis de datos espacio-temporales en áreas como la Epidemiología, Agricultura y Medicina.
Improving the transparency of randomised trial manuscripts: results from a randomised trial to enhance author adherence to CONSORT guidelines

David Blanco\textsuperscript{1,2}, Sara Schroter\textsuperscript{3}, Adrian Aldcroft\textsuperscript{3}, David Moher\textsuperscript{4}, Isabelle Boutron\textsuperscript{5}, Jamie J Kirkham\textsuperscript{6} and Erik Cobo\textsuperscript{1}

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\textsuperscript{6}Department of Biostatistics, University of Liverpool, Liverpool, UK.

Improving the transparency and accuracy of randomised controlled trials (RCT) is key to reduce waste in biomedical research. The Consolidated Standards of Reporting Trials (CONSORT) Statement (1) and other reporting guidelines help authors report research methods and findings in a way that they can be understood by a reader, replicated by a researcher, used by a doctor to make a clinical decision, and included in a systematic review (2).

In recent years, biomedical journals have taken different actions to boost the completeness of reporting of published randomised trials. One of the most popular ones consists of requiring authors to submit a populated CONSORT checklist together with their manuscript indicating page numbers corresponding to each item. However, journals usually lack further actions throughout the editorial process to ensure that the corresponding information to each item is actually reported in the manuscript. This has been hypothesized to be one of the reasons why this strategy has not yet achieved optimal results.

In an effort to take full advantage of the submission of populated CONSORT checklists, we are currently performing an RCT (3) in collaboration with BMJ Open, an open-access journal that publishes medical research. Our goal is to evaluate the impact of the following actions on the transparency of the randomised trials published in BMJ Open: 1) assessing during peer review the consistency between the CONSORT checklist submitted by authors and the information that is reported in the manuscript, and 2) writing up a peer review report providing feedback to authors on the inconsistencies found.

In this presentation, we will present the preliminary results of our RCT. Moreover, we will detail how we intervened the editorial process, how authors reacted to the intervention and what challenges we faced. Finally, we will briefly describe other interventions that could be implemented throughout the editorial process in order to improve author adherence to CONSORT and other reporting guidelines.
We hope this presentation will raise awareness on the importance of reporting research transparently and will make attendants reflect on the vital role of biomedical journals taking a step forward to ensure that this happens.

**Keywords:** Reporting guidelines, transparency, adherence, randomised trial.

## References


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**Evaluation of the prognosis of HPV negative but p16 positive oropharyngeal cancer patients in an international collaborative study (EPIC-OPC)**

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**Background:** In 2007, the International Agency of Research on Cancer concluded that there was sufficient evidence for the carcinogenic role of HPV16 in oropharyngeal cancers (OPCs) [1]. However, this cancer etiology is multifactorial and tobacco and alcohol consumption also play an important role. HPV-DNA detection has proven to be insufficient to attribute causality, thus it is crucial to incorporate additional biomarkers to assess the biological and oncogenic potential of HPV, such as p16 expression [2]. In addition, it has been shown that patients with double positivity for HPV-DNA/p16 have a better overall survival compared to any other combination [3]. In spite of this, some scientific societies recommend only performing p16 determination for HPV-relatedness classification. In order to clearly describe the characteristics of HPV-negative/p16-positive patients, since most cohorts had amassed relatively few cases, an international consortium has been created.

**Objectives:** (1) To describe the distribution of the double HPV-DNA and p16 detection in two retrospective cohorts (Spain and Germany); (2) To identify the
detectors for the combinations of the double detection of HPV-DNA and p16 in OPC patients.

**Methodology:** Information on sociodemographic and clinical characteristics at diagnosis of primary OPC from patients from two different centers was recruited. Pairwise tests of independence for nominal data were calculated. In order to determine the factors associated to HPV DNA infection and p16 overexpression, multinomial logistic regression analysis was performed using both Frequentist and Bayesian approaches [4, 5].

**Results:** A total of 1563 patients with a primary OPC were recruited in the two cohorts between 1991 and 2018. These cases were distributed according to their combination of the HPV and p16 detection as follows: double HPV/p16-negative (n=1163), HPV-negative/p16-positive (n=67), HPV-positive/p16-negative (n=71) and double HPV/p16-positive (n=262). There were significant differences between double negative and double positive cases for all the variables; and between double positive and HPV/p16 discordant results for some variables such as recruiting centre, tumour subsite, tumour staging, smoking status, alcohol consumption, period of diagnosis. In tonsil and base of the tongue sites, the risk factors for being double positive were: young cohorts, advanced tumour stage, non-smokers, non-drinkers and diagnosed in recent years. In other oropharyngeal sites, the risk factors to be double positive were to be a non-smoker and a non-drinker patient. The adjusted OR from the Frequentist and the Bayesian, including a random effect on the intercept with center as a grouping variable, approaches were similar, being the magnitudes of the second one slightly larger.

**Conclusions:** We observed a clear difference in the factors associated to double negative and double positive patients by subsite. HPV-negative/p16-positive patients shared the same determinants with the double negative patients. HPV-positive/p16-negative patients group resembles more to the double negative than to the double positive group confirming that both determinations are necessary to correctly classify patients with OPC. Although we have not observed any difference between the results from the Frequentist and the Bayesian approaches, we propose to use the second one to better characterize this patients.

**Keywords:** Oropharyngeal Cancer (OPC), Human Papillomavirus (HPV), Odds Ratio (OR), Multinomial Logistic Regression, MCMCglmm

**References**


Mena M, Taberna M, et al. (2018). Double positivity for HPV-DNA/p16INK4a is the biomarker with strongest diagnostic accuracy and prognostic value for
The human microbiota is the collective genomes of the microorganisms that live inside and on the human body. It is composed by many types of microbes as bacteria, archaea, fungi, protists and viruses. These microorganisms resides either in or on of a number of human tissues and biofluids, including the skin, mammary glands, placenta, seminal fluid, uterus, ovarian follicles, lung, saliva, oral mucosa, conjunctiva, biliary and gastrointestinal tracts. Microbiota have been found to be crucial for the health of their host. The human microbiota and their interactions with the host play an important role in basic biological processes and in the development and progression of major human diseases such as infectious diseases, liver diseases, gastrointestinal cancers, metabolic diseases, respiratory diseases, mental or psychological diseases, and autoimmune diseases.

Note that microbiota is inherently dynamics. Thus, longitudinal microbiota analysis can provide rich information of short-and long-term trends of microbial communities. In addition, a common way to express the microbiota data is using relative proportions. As a consequence, compositional data analysis is a valid approach to analyze the microbiota sequencing data.

In this work, we propose a dynamical model in order to analyze trends of microbiota timeseries with a Dirichlet response variable. We will present different modeling proposals and estimation methods.

**Keywords:** Microbiota, Multivariate Time Series, Dynamic Models, Dirichlet, Bayesian inference, Compositional Data.
Mediation analysis and Network Mendelian Randomization: methods for the estimation of the causal effect

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\textsuperscript{3}Universidad Complutense de Madrid (UCM), Madrid, España

Introduction: Causal Mediation Analysis (CMA) is a statistical method that allows investigators to consider possible mediator variables in the causal inference chain. The use of genetic variants as Instrumental Variables (IV) in causal inference studies is increasing in the last years. IVs usage in the CMA context is known as Network Mendelian Randomization (MR), but there are few studies describing the appropriate methodology. Classic CMA approaches, such as Potential Outcome Framework (POF), i.e., the counterfactual approach, and Structural Equation Models (SEM), are suitable to implement IVs.

Objective: To compare several CAM approaches including POF, SEMs and Network MR using simulated and real data, within a study on the causal mediation effect between subtypes of T2DM (new-onset NODM and long-standing LSDM) and obesity in pancreatic cancer aetiology.

Methods: Study population (real data): We considered a European multicentre case-control study, the PanGenEU study, with 2,108 PC cases and 1,540 controls. The majority had genotype data. Simulation study: We simulated the outcome variable, as well as the exposure, the mediator and the IVs. The outcome and mediator were considered both as continuous and as binary variables. The exposure was considered in all analyses as a binary variable. We first considered a classic mediation analysis approach using subtypes of T2DM as exposure, obesity as mediator, and vice versa, and pancreatic cancer as outcome using the POF and SEMs. To implement IVs within these methods, we defined specific IVs for the mediator and exposure variables. These IVs were considered as a genetic score of T2DM or obesity-related genetic variants, and included in the fitted final model. In the case of Network MR, we implemented IV methods such as the Wald Ratio (WR) and Multiple-Stage-Least-Squares (MSLS).

Results: We obtained similar results in terms of Odds Ratios (OR) and 95\% Confidence Interval (CI) when comparing the classical mediation analysis approaches and their counterparts using IVs. For instance, in POF, the total causal effect of obesity as mediator in the causal pathway between LSDM and PC risk was 1.68 (95\%CI: 1.37-2.06) and 1.58 (95\%CI: 1.16-2.15), respectively. Regarding Network MR considering the IV methods (WR and MSLS) for binary variables, results were not consistent and need further development.

Conclusions: To obtain the correct causal estimation effect in CMA, it is important to consider the proper methodology. The implementation of IVs in the counterfactual and SEM was possible but requires further development in the context of Network Mendelian Randomization.
Session 8: Spatial statistics and statistical methods in agriculture  
September 6th, 16:30 to 17:45  
Chair: Elena Lázaro

RAAPS: A Shiny App for Risk Assessment around Pollution Sources

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La importancia de evaluar los riesgos de la polución alrededor de fuentes de emisión de gases es un importante problema espacial estudiado en epidemiología. La metodología usada de casos y controles, dan una versatilidad para el análisis en este tipo de estudios. Como ejemplo, cuando evaluamos el impacto de la polución emitida por industrias, es habitual pensar en la distancia al foco como factor de riesgo en el modelo, así, cuando la localización de los casos y controles son conocidos por el investigador, se pueden proponer un número de modelos que evalúen el impacto de dichas fuentes de polución en la distribución espacial de los casos teniendo en cuenta la distribución de los controles. Sin embargo, este tipo de análisis requieren una manipulación compleja de los datos tal como mapas, visualización espacial, estadísticas espaciales entre otros. También, RAAPS representa una oportunidad y estímulo para quienes generan datos. El personal de centros como hospitales podría trabajar directamente sus datos, optimizando el poco tiempo y recursos que disponen para tareas investigadoras.

Por este motivo, hemos desarrollado una App en el entorno Shiny llamada RAAPS, del inglés (Risk Assessment App around Pollution Sources) con la intención de ayudar a los investigadores que utilizan el método de casos y controles en el campo de la salud. La App se compone de una simple interface donde cargamos los datos, así como la localización de las fuentes de riesgo (en nuestro caso industrias), en una región de estudio fácil de crear y reconocer. De manera similar, un análisis estadístico evalúa el impacto de la localización de las industrias sobre los casos hallados de enfermedad. Esto daría lugar a una importante herramienta a hospitales para la vigilancia epidemiológica y la monitorización de riesgos. El uso de esta App se ilustra con un conjunto de datos con las localizaciones de diferentes tipos de cáncer (pulmón, estómago y riñón) en la ciudad de Alcalá de Henares (Madrid, España).

Se usaron varias industrias cerca de la ciudad para el análisis.
Spatial modelling of *Xylella fastidiosa* distribution: two case studies in the Mediterranean Basin

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*Xylella fastidiosa* (*Xf*) is a phytopathogenic bacterium with a wide range of host plants, characterized by its genetic diversity. In this study, the effects of climatic and spatial factors on the distribution of *Xf* in two areas of the Mediterranean Basin were analyzed: Apulia (Italy) and Alicante (Spain). The pathogen was confined to the American continent and Taiwan, but in 2013 it was detected in Europe, in Apulia region, where extensive areas were affected by the olive quick decline syndrome, caused by *Xf* subsp. *pauca*. Almond leaf scorch, caused by *Xf* subsp. *multiplex*, was detected in 2017 in Alicante (Spain). Presence/absence data of *Xf* in the official surveys in both regions, Apulia and Alicante, were analyzed using a Bayesian approach through the Integrated Nested Laplace Approximation (INLA) methodology (Rue et al., 2009). Climatic covariates were obtained from the WorldClim database (Fick and Hijmans, 2017). In addition, a categorical variable was included according to Purcell’s levels of disease severity based on minimum winter temperature from < 1.1°C to > 4.5°C. These thresholds were defined in North America for Pierce’s disease of grapevine, caused by *Xf* subsp. *fastidiosa*, and it is not known if those thresholds can be extrapolated to other *Xf* subspecies or geographic regions. Due to the different surveillance and sampling strategies carried out in the two study areas, the available georeferenced data presented a different structure. In Apulia, data were observed at continuous locations occurring within a defined spatial domain (geostatistical data). Therefore, the spatial effect was included using the Matérn covariance function, approximated as a solution to a Stochastic Partial Diferential Equation (SPDE) (Lindgren et al., 2011). In Alicante, the samples were overlapped on a grid, so they were considered lattice data and the spatial effect was included through a conditional autoregressive structure (iCAR). Results showed that in Alicante the *Xf* was detected in all four Purcell’s levels, illustrating the climatic adaptability of *Xf* subsp. *multiplex*. In Apulia, with much lower winter temperature variability than Alicante, only the high risk Purcell’s class was observed. In both regions, the spatial structure had a strong influence in the models, but not the climatic covariates. In conclusion, disease spread was largely defined by the spatial relationship between geographic locations.

**Keywords:** Hierarchical Bayesian models, INLA, *Xylella fastidiosa*.

**References**


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A procedure for Satellite image calibration using UAV imagery for agriculture applications

Amal Chakhar¹, David Hernandez-Lopez¹ and Miguel A. Moreno¹,²

¹Institute for Regional Development (IDR), University of Castilla-La Mancha, Albacete, Spain.

The launch of Sentinel 2 (A and B) satellites has boosted the development of many applications for instance describing crop status and variability. Sentinel 2 offers ground sample distance (GSD) equal to 10 m and combined constellation revisit time equal to 5 days (ESA, 2015). According to (Padrò et al., 2018), the made efforts in combining different platforms products still insufficient because there is a high risk of uncertainty that these product receive an inadequate radiometric correction that can not allow sensors to perform in coordinated way. The radiometric correction is applied to spaceborne or airborne imagery with the purpose to extract physical earth surface parameters for example reflectance quantities (Richter and Schläffer, 2016). Based on its many features, such as the flexibility along with the high spectral, temporal and spatial resolution (Herrero-Huerta et al., 2014) imagery obtained from multispectral sensors onboard Unmanned Aerial Vehicles (UAV) is expected to complement field data. In this study, a detailed analysis and comparison of multispectral imagery provided by Sentinel 2 and UAV was performed. The Sentinel 2 and the resampled UAV products was evaluated by considering, first, the intercomparison of NIR and Red reflectance bands, and second, by the intercomparison between the NDVI of the two products at the level of the whole flight zone as well as at different crop level. When downsampling the UAV imagery to the Sentinel 2 pixel size we tested two methods: the nearest neighbor and the average of the UAV pixel values. Likewise, we aimed the elaboration of models with the purpose to understand the relationship between the two different sensor platforms in the study area, and to develop a predictive model for barley, vineyard, almond and olive. Based on multitemporal band-by-band comparison, we concluded that the average resampling method provide better correlation results and lower RMSE than the Nearest Neighbor method. The NDVI fitting models at the whole flight zone suggest that at least 67% (which corresponds to the day with the lowest R2) of the variation in the data can be explained by the linear model. Concerning the predictive models, for barley, we obtained good linear regression model with
strong correlation coefficient varying from 0.86 in 2018.01.19 to 0.921 in 2018.06.13. For almond, vineyard and olive we did not obtain a good linear regression models as the barley model, therefore, we concluded that the herbaceous provide better results than the ligneous.

**Keywords:** Sentinel 2, UAV, reflectance bands, NDVI, calibration, resampling, downsampling, predictive models.

**References**


**Application of laser scanner systems in precision viticulture**

Ana del-Campo-Sanchez¹, Miguel Moreno¹, Rocio Ballesteros¹ and David Hernandez¹

¹AgroForestry and Cartographic Precision Research Group, University of Castilla – La Mancha, Spain

Three-dimensional (3D) modelling is a growing remote sensing technique for the agroforestry sector (i.e.[1]). 3D scanning by laser scanner allows to collect most of geometries from agroforestry scenarios using only one geomatic product, termed point clouds. This massive digitization of spatial data involves a wide range of information to be process by powerful computing tools. These tools make possible to extract useful information in the decision making process for management and conservation of agricultural and forestry environments. Scientific community is making much efforts to improve these methodologies and tools as they decrease costs and increase precision and density points.

However, the main challenge of this technology is to close hull points properly to define scanned surfaces. Point clouds captured by laser scanning systems only provide isolated data of the intercepted surfaces by light pulses emitted from the sensor. Therefore, there is no spatial data between one single point and its neighbours. For that reason, defining scanned surfaces relating one point to the next one involves a high computational complexity [2], primarily in complex objects such as the ones in agroforestry systems. Thus, the current challenge consists of moving from a model...
expressed as a point cloud to a conventional three-dimensional model that allows us to extract metric characteristics that provide information of interest in the process of analysis or characterization of the modelled object.

Within the framework of precision viticulture, determining the vigour of the vines that composes the vineyard permits take decisions about the inputs to allocate in the plot, determining possible differences in harvest and the probability of diseases and pest affections. Thus, determining the actual volume occupied by each vine could derive a much precise estimation of vigour of the vines.

For these purposes, this research work presents a strategy for determining the volume of individual vines from point clouds collected with a static terrestrial laser scanner (TLS) and an integrated mapping mobile system (MMS), using several algorithms implemented and adjusted for each case (such as convex hull, OctoMap and the L1-medial skeleton method [3–5]), which constitute a problem in the context of biometrics.

Results shows that the two first methods are appropriate when using TLS systems, but they are totally unable for low point cloud density. This problem is solve using the skeleton calculation. These results mean a great advance in the use of laser systems in agroforestry applications, showing the capability of these methods of extracting useful information from dense and sparse point cloud products.

**Keywords:** Point cloud, Mapping Mobile System, Terrestrial Laser Scanner, Agroforestry Precision.

**References**


Water stress and yield assessment for vineyard orchard by using multispectral and RGB very high resolution aerial images

P. López-García¹, Moreno, M.A.¹, J.F. Ortega¹ and Ballesteros, R.¹

¹Institute of Regional Development, Campus de Albacete

This study was developed in the framework of precision viticulture and very high resolution remote sensing by using drones. The experiment was carried out along season 2018 in Fuente-Álamo (Albacete), in one experimental plot of 7.5 ha. The studied crop was vineyard (Vitis vinifera L.). Six different irrigation treatments were applied in order to study plant behavior regarding water stress and yield. The treatments applied were: 1) rainfed; 2) irrigation; 3) and 4) application of irrigation water including two types of salts; 5) and 6) same salty irrigation treatments only from veraison. 1000 m³/ha were applied to all irrigation treatments. Six flights were made with a UAV with RGB and multispectral cameras on board. The acquired images were photogrammetrically processed to obtain, among other geomatic products, an orthoimage in the RGB range and the multispectral range for each flight event.

These orthoimages were the starting point for calculating some of the main vegetation indexes (VIs) traditionally used for vegetation characterization. Linear relationships between these VIs and accumulated water stress and yield were analyzed. Furthermore, relationships between different multispectral and visible bands were studied using artificial neural networks (ANNs).

The study of water potential did not show significant differences between the different treatments until veraison. There were no differences between treatments with different types of applied salts, neither there were significant differences between the treatments with respect to yield. However, differences were evident between irrigation and rainfed treatments.

Statistical analysis of multispectral VIs and linear relationship with water stress along growing cycle showed Renormalized Difference Vegetation Index (RDVI) as the index with the best adjustment ($R^2 = 0.724$, RMSE = 6.564), while for yield the studied VIs showed inadequate adjustments ($R^2 < 0.5$). In the case of RGB-based VIs, the adjustment for linear relationships and water stress showed the most accurate adjustments for Excess Red Vegetation Index (ExR) ($R^2 = 0.715$, RMSE = 3.299 MPa-days). Adjustment for linear models between yield and RGB VIs was not significant, as it did for multispectral VIs. The use of ANNs, both of water stress and yield, for the two spectral ranges showed a significant improvement of adjustments, being remarkable for ANNs and RGB bands with adjustments of $R^2 = 0.985$, RMSE = 0.313 MPa-days for stress determination and $R^2 = 0.982$, RMSE = 0.216 kg/vine for yield. Thus, networks can be considered the best alternative for the description of these highly non-linear relationships.

Keywords: multispectral range, RGB range, UAVs, Vitis vinifera L., water stress, yield
# Time schedule

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<td>Sesión 1: Medical studies I</td>
<td>Sesión 5: Multivariate analysis</td>
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<td>Sesión 2: Statistical ecology</td>
<td>Sesión 6: Prediction models</td>
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Thursday, September 5th

08:30 Registration

09:00 Welcome

SESSION 1: MEDICAL STUDIES I
Chair: Lore Zumeta Olaskoaga

09:30 “Development of statistical methodologies to work with ELISpot assays”, Yovaninna Alarcón Soto.


10:00 “Modelos multinivel aplicados a la comparación de características y resultados de pacientes con diagnóstico de cáncer colorrectal según su vía de detección”, Sergio Moreno.

10:15 “A statistical model for analysing the effects of treatments on multivariate time series of microbiota data”, Irene Creus Martí.

SESSION 2: STATISTICAL ECOLOGY
Chair: Joaquín Martínez Minaya

10:30 “Big problems, integrated solutions”, Blanca Sarzo.

10:45 “Mixed models for exploring sardine fishing in the Mediterranean Sea”, Gabriel Calvo.

11:00 “From data-limited to data-rich: the evolution of sardine in the Bay of Biscay”, Leire Citores.

11:15 “Variable selection for genome wide association analysis in plant breeding”, Iker Oyanguren.

11:30 Coffee break

12:00 R Course

14:00 Lunch

SESSION 3: SURVIVAL ANALYSIS AND OPTIMUM EXPERIMENTAL DESIGN
Chair: Yovaninna Alarcón Soto

15:30 “Prognostic factors in the survival of women with breast cancer through a Bayesian competitive risk model”, Sandra Ramírez.
15:45 “Evaluación basada en la modificación de la historia natural de la efectividad de un programa poblacional de atención a la insuficiencia cardíaca”, Ania Goroztiza.

16:00 “Statistical Modeling in sports injury prevention”, Lore Zumeta Olaskoaga.

16:15 “Pensar antes de actuar: introducción al diseño óptimo de experimentos”, Sergio Pozuelo Campus & Irene García Camacha.

16:45 Coffee break

SESSION 4: BIOSTATNET
Chair: Irene García Camacha

17:00 “Three-level hierarchical modelling of longitudinal data: Application to the study of worldwide obesity prevalence”, Diana Marcela Pérez Valencia.


17:30 “CompARE: a web app to study composite endpoints”, Marta Bofill Roig.

17:45 “Fitting Bayesian Dirichlet regression using the integrated nested Laplace approximation”, Joaquín Martínez Minaya.

18:00 “Spatial conditional overdispersion models: Modelling mothers’ postnatal screening period”, Mabel Morales Otero.

POSTER SESSION
(18:15-19:15)


5. “Robustness of design for accelerated failure time models with Type I censoring”, Irene García-Camacha Gutiérrez


19:30 Social and tourist activity

22:00 Dinner
Friday, September 6th

SESSION 5: MULTIVARIATE ANALYSIS
Chair: Sergio Pozuelo Campos

09:30 “The smart potential of European regions”, Belmonte Leticia Cortés.

09:45 “Aplicación de la Teoría de Respuesta al ítem para armonizar diferentes escalas de depresión en el proyecto ATHLOS”, Laia Egea Cortés.

10:00 “Caracterización del dimorfismo sexual a nivel craneal mediante análisis discriminantes”, Luis Sánchez Martínez.

10:15 “From high-dimensional to functional data: application in SNPs data.”, Harold Hernández Roig.

SESSION 6: PREDICTION MODELS
Chair: Blanca Sarzo

10:30 “Overview and new proposals in the development of prediction models with complex survey data”, Amaia Iparraguirre.

10:45 “Estudio comparativo de métodos de estimación de los parámetros del modelo de regresión logística con datos de encuestas con diseño complejo”, Ane Villanueva.

11:00 “Targeted lipidomic profiling reveals a lipid mediator signature specific of acute-on-chronic liver failure”, Arantxa Urdangarin Iztueta.


11:30 Coffee break

MESA REDONDA: “EL PAPEL DE LAS MUJERES EN LA ESTADÍSTICA”
(12:00-13:30)

13:30 Lunch

SESSION 7: MEDICAL STUDIES II
Chair: Guillermo Villacampa

15:00 “Improving the transparency of randomised trial manuscripts: results from a randomised trial to enhance author adherence to CONSORT guidelines”, David Blanco.
15:15 “Evaluation of the prognosis of HPV negative but p16 positive oropharyngeal cancer patients in an international collaborative study (EPIC-OPC)”, Sorina Sferle.


16:00 Break

SESSION 8: SPATIAL STATISTICS AND STATISTICAL METHODS IN AGRICULTURE
Chair: Elena Lázaro

16:30 “RAAPS: A Shiny App for Risk Assessment around Pollution Sources”, José Luis Gutiérrez Espinosa.


17:00 “A procedure for Satellite image calibration using UAV imagery for agriculture applications”, Amal Chakhar.


17:30 “Water stress and yield assessment for vineyard orchard by using multispectral and RGB very high resolution aerial images”, Patricia López.
Location

Most of the congress will be in the auditorium (Salón de actos). The course will be in classroom 0.1 and meals in the cafeteria.
# List of participants

In alphabetical order:

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<th>Nr.</th>
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