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# The use of Meta-Regression Analysis to harmonize LCA literature: an application to GHG emissions of 2<sup>nd</sup> and 3<sup>rd</sup> generation biofuels

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## Abstract

This article presents the results of a literature review performed with a *meta-regression analysis (MRA)* that focuses on the estimates of advanced biofuel Greenhouse Gas (GHG) emissions assessed with a Life Cycle Assessment (LCA) approach. The mean GHG emissions of both second (G2) and third generation (G3) biofuels and the effects of factors influencing these estimates are identified and quantified by means of specific statistical methods. 47 LCA studies are included in the database, providing 593 estimates. Each study estimate of the database is characterized by *i*) technical data/characteristics, *ii*) author's methodological choices and *iii*) typology of the study under consideration. The database is composed of both the vector of these estimates – expressed in grams of CO<sub>2</sub> equivalent per MJ of biofuel (g CO<sub>2</sub>eq/MJ) – and a matrix containing vectors of predictor variables which can be continuous or dummy variables. The former is the dependent variable while the latter corresponds to the explanatory variables of the meta-regression model. Parameters are estimated by means of econometrics methods.

Our results clearly highlight a hierarchy between G3 and G2 biofuels: life cycle GHG emissions of G3 biofuels are statistically higher than those of Ethanol which, in turn, are superior to those of BtL. Moreover, this article finds empirical support for many of the hypotheses formulated in narrative literature surveys concerning potential factors which may explain estimates variations. Finally, the *MRA* results are used to address the harmonization issue in the field of advanced biofuels GHG emissions thanks to the technique of *benefits transfer using meta-regression models*. The range of values hence obtained appears to be lower than the fossil fuel reference (about 83.8 in g CO<sub>2</sub>eq/ MJ). However, only Ethanol and BtL do comply with the GHG emission reduction thresholds for biofuels defined in both the American and European directives.

**Keywords:** Biofuels, GHG, LCA, Meta-analysis

# 1 Introduction

This article addresses the environmental evaluation issues of advanced biofuels. It focuses on a specific environmental evaluation method – Life Cycle Assessment (LCA) – and its estimates of second (G2) and third generation (G3) biofuels greenhouse gas (GHG) emissions. The mean Global Warming impact indicator, expressed in grams of CO<sub>2</sub> equivalent per MJ of biofuel (g CO<sub>2</sub>eq/MJ), and the effects of factors influencing these estimates are characterized and quantified using a *meta-regression analysis (MRA)*: a quantitative research method to review and synthesize empirical literature. This research is of primary importance as this measure may be interpreted as an estimate of the contribution to climate change of advanced biofuels.

First generation (G1) biofuels have been developed to provide a substitute for fossil fuels in order to enhance energy independence and mitigate climate change. They mainly correspond to Ethanol and biodiesel produced from conventional crops such as sugar cane, sugar beet, wheat, corn, rapeseed, sunflower, etc. Nevertheless, G1 biofuels have come up against sustainability issues. Indeed, the use of agricultural commodities for the production of biofuels induces an additional demand for these crops and, consequently, an increased use of arable land. Furthermore, it has been suggested that it may induce a rise of food prices [1]. Additionally, many life-cycle based studies point out that G1 biofuels do not reduce GHG emissions as significantly as expected [2]. As a consequence, G2 and G3 biofuels (referred to in this paper as advanced biofuels) from biomass residues, non-alimentary crops and wastes have been developed in the recent years. These biofuels seem to be more efficient than G1 biofuels in terms of land use, food security, GHG emission reductions and other environmental aspects [3].

G2 Ethanol is obtained from the biochemical conversion of lignocellulosic biomass<sup>1</sup> and synthetic diesel from biomass, also known as BtL (Biomass to Liquids) or biomass FT-diesel, is produced by the thermochemical conversion of lignocellulosic biomass. G3 biofuels are produced from microalgae using algal oil for biodiesel production from conventional transesterification (a.k.a Fatty Acid Methyl Ester, FAME) or hydrotreated vegetable oil (HVO) (See Appendix A for further details). Advanced biofuels are currently either in research and development or demonstration phase and still need further improvements to be commercially viable.

Some states have set ambitious production targets for biofuels, supported by subsidies and legislative incentives. In the European Union (EU), the Renewable Energy Directive (RED, [4]) requires the use of 10% of renewable energies in the transport sector by 2020 (in 2009, the share was 3.6%). To achieve this goal, the contribution of biofuels produced from lignocellulosic materials, wastes and residues is considered to be twice that made by other biofuels. This can be viewed as an incentive to develop advanced biofuels. In the United States (US), the Renewable Fuel Standard (RFS2, [5]), under the US Energy Independence and Security Act of 2007, requires the use of 136 billion liters of biofuels by 2022 (in 2009, 41.9 billion liters were mandated). It specifies that 79,3 billion liters must be of "advanced biofuels" and "cellulosic biofuels" (the definition of "advanced biofuels" in the RFS2 is different from the one adopted in this paper and will be clarified later on). In addition, other countries (Australia, China, Japan, New Zealand, Brazil and others) have already been actively developing next generation biofuels and feedstock although there is little policy support in these regions [6].

Furthermore, the EU and the US set a list of sustainability requirements for biofuel production. In both regions, the only mandatory quantitative criterion is related to life cycle GHG emissions calculated using the LCA method. The RED sets minimum life cycle GHG emission savings for all biofuels compared to a fossil fuel reference. These savings are of 35% since 2009, and will be of 50% in 2017 and 60% from 2018 for

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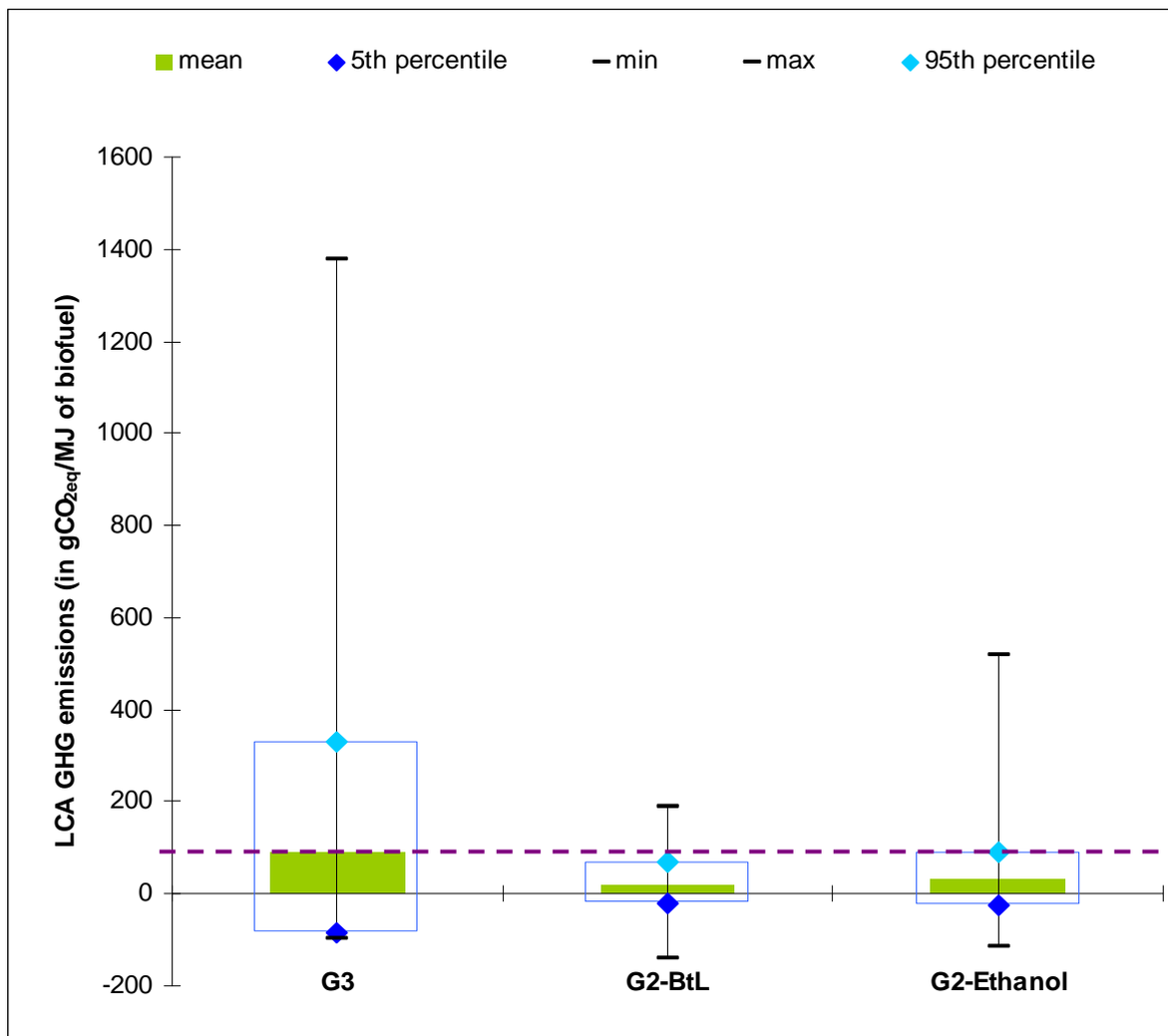
<sup>1</sup> Lignocellulosic biomass refers to annual crop residues (e.g. corn stover), forest residues, herbaceous energy crops (e.g. switchgrass, miscanthus) and woody biomass (e.g. poplar, eucalyptus).

new biofuel plants. The RFS2 also sets minimum life cycle GHG emission savings that biofuels have to comply with in order to be eligible for appropriate subsidies. Those savings are set to 20% for first generation biofuels, 50% to be considered as "advanced biofuel" (as defined in the RFS2, different from our definition) and 60% to be considered as "cellulosic biofuel".

Those GHG emission requirements as well as biofuel incorporation targets are clearly in favour of G2 and G3 biofuels. This shows the will of policy makers to support their future development compared to G1 biofuel. That is one of the reasons why we choose to focus on advanced biofuels in this study.

We choose to conduct our literature analysis by reviewing only LCA studies assessing Global Warming impact indicators, *i.e.* GHG emissions, for the following two reasons. First, one of the main objectives for developing biofuels is to reduce global GHG emissions in order to mitigate climate change. As an illustration, recall that the only quantitative mandatory requirement for biofuel sustainability is related to life cycle GHG emission savings in the EU and in the US. Thus, it appears important to check advanced biofuel compliance with this requirement by comparing their life cycle GHG emissions with those of a fossil fuel reference. Second, a significant literature already exists that assesses GHG emissions of advanced biofuels using the LCA approach. Hence a sufficient number of studies is available to investigate this issue. Note that because GHG emissions have an environmental impact at a global scale (GHG emission effects do not depend on the place where they have been emitted), this literature review includes worldwide studies.

The first applications of LCA to biofuels to measure a Global Warming impact indicator were carried out on G1 biofuels in the 90's (such as Kaltschmitt *et al.* [7]). Since, numerous LCA studies were conducted to analyze G2 and G3 biofuel pathways. Despite this substantial literature, the extent to which advanced biofuels may have lower GHG emissions than the fossil reference remains a subject of debate. While the majority of these studies shows GHG benefits for advanced biofuels compared to a fossil fuel reference, some authors come to the opposite conclusion. For instance, LCA GHG emission results selected for this study (47 studies providing 593 GHG emission results, see next section for more details) range from -142 (G2) to 1378 (G3) g CO<sub>2</sub>eq/MJ of biofuel (see Figure 1); the greatest variability of GHG emission results being for G3 biofuels.



**Figure 1 - GHG emissions extrema for bibliographic results of G2 and G3 biofuel LCA studies (47 studies, 593 observations)**

When looking at Figure 1, one can wonder *i)* if there is a consensus about GHG emission benefits from advanced biofuels and *ii)* why there is so much variation among results of these studies even though they are all investigating the same phenomenon.

Actually, even if the LCA approach is consistent throughout, each study – by nature – concerns different pathways and uses specific data and methodological assumptions. Previous narrative surveys of biofuel LCA studies mention that LCA results are inconclusive regarding GHG emission performances of advanced biofuels [8–13]. According to these literature reviews, LCA GHG emission results for advanced biofuels vary significantly depending on various factors such as: the assumptions made to describe the biomass production step (model used to estimate N<sub>2</sub>O emissions and inclusion of direct and indirect land use change), the data used to describe the biomass conversion into biofuel and the general LCA methodological choices (system boundaries, the method used to account for coproducts impacts, etc.). While these indicative results from literature reviews are really useful, primary study results remain difficult to compare because of differences in technical data or methodological choices.

As a consequence, it is quite difficult to attempt any summary and to form an accurate opinion on this topic using classical literature reviews methods. In particular, it seems hard to provide one GHG emission estimate appropriate for advanced biofuels.

Since most studies are inconclusive, their results may not be relevant for decision support [14]. There is a strong need for harmonization of LCA results, especially for policy makers or investors, as suggested by Heath and Mann [15] with the "*LCA harmonization project*". The purpose of harmonization, as defined by

Heath and Mann, is to identify and quantify key factors that influence the environmental impacts for a technology or product in order to be more conclusive concerning its real environmental performances. At present, few studies have tried to harmonize GHG emission results from various LCA studies for advanced biofuels. For instance, Handler *et al.* and Liu *et al.* [16,17] propose to harmonize GHG emission results for G3 biofuels by normalizing their LCA models using same methodological assumptions and same generic pathways.

Although it is not possible to calculate one GHG emission estimate appropriate for all advanced biofuels, we believe it remains possible to determine central tendencies based on the distribution of previous study results. To do so, this article proposes an alternative summary to previous literature reviews, using the *meta-analysis (MA)* methodology to describe and synthesize existing estimates of the LCA GHG emissions of advanced biofuels.

*MA* is a quantitative research method developed to compare and/or combine outcomes of different individual quantitative studies, named primary studies, with more or less similar characteristics that can be controlled for [18]. By nature, each result from a primary study (called an estimate) may be quoted to illustrate the uncertainty of estimates. Estimates of previous studies are grouped together in a database, called meta-database, according to one or more differentiating characteristics. These estimates become then the observations, also named *effect-size (e-s)* (see below), of the meta-database whereas the differentiating characteristics become their potential explicative variables. In a *MA* framework, the *e-s* is assumed to be a function of these explicative variables; function which can be specified and assessed. When this *meta-function* is estimated by the means of multi-regression techniques, *i.e.* specific econometrics estimators, the *MA* is called a *meta-regression analysis (MRA)*<sup>2</sup>. This multivariate setup allowed by the meta-regression framework is very useful in the field of literature reviews as it enables us to statistically identify and quantify – *ceteris paribus* – the effect of the most influential characteristics on the *e-s*. Thus, compared to narrative literature reviews, the *MRA* methodology – thanks to its multivariate setup – gives the opportunity to test the influence of specific characteristics, after having controlled for the effect of other ones. Besides, a "*meta-regression*" framework allows to produce an estimation of the mean *e-s* weighted by the systematic influence of its main drivers. Indeed, once statistically estimated, the meta-function can be used to deduce original values of the *e-s* by specifying new values for the main drivers identified corresponding to relevant case studies. This technique of *benefits transfer using meta-regression models*, as it is named in the *MA* literature, may be a particularly well adapted methodology to deal with the so-called harmonization issue specific to the LCA literature.

The literature of LCA studies estimating advanced biofuels GHG emissions is now large enough to support a statistical assessment of this measure of the mean Global Warming impact indicator. The primary purpose of this *MRA* is to identify and quantify by statistical estimates which factors among *i)* technical data/characteristics, *ii)* author's methodological choices and *iii)* typology of the study under consideration have an impact on variations of the GHG emission estimates. The second purpose of this *MRA* is to generate a distribution of the potential GHG emissions of advanced biofuels and to characterize the mean Global Warming impact indicator and its standard deviation across G2 and G3 biofuels. We investigate – through an application – the potential for *MRA* to synthesize LCA literature by highlighting the main determinants of result variability in order to perform harmonization.

This paper is organized as follows. Section 2 is a brief summary of both LCA approach applied to biofuels and *MRA* methodology. Section 3 is a description of the meta-database in which the *e-s* and explanatory variables are described. Meta-regression models and the associated results are presented and analyzed in Section 4. Main conclusions and methodological discussion are presented in Section 5.

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<sup>2</sup> So defined, *MRA* may be viewed as a subset of *MA* in the literature.

## 2 Methods

First, this section briefly presents the LCA approach and then summarizes how it has been used in the literature to estimate Global warming impact indicators of advanced biofuels. Second, the meta-regression methodology is briefly presented. Both sections enable a better understanding of the *e-s* and explanatory variables of the *MA*.

### 2.1 General presentation of LCA method

Life Cycle Assessment (LCA) is a method based on ISO standards 14040/14044 [19,20] aimed at assessing several potential environmental impacts of a product or a service during all of its life cycle. This approach takes into account all steps of a product's life cycle: from the extraction of natural resources necessary for its production (oil, coal, gas, etc) to its end of life or destruction ("Cradle to Grave" analysis). The LCA approach enables the characterization of potential environmental performances of a production system in order to identify potential improvements and is a relevant tool for decision makers.

The methodological framework for LCA set by international ISO standards is divided into 4 steps:

1. Goals and Scope of the study: This step deals with the definition of questions that we want to answer in the study and the final users of the results. Hence all methodological assumptions, i.e. the scope of the study (system boundaries, functional unit, method to account for coproducts, environmental impact indicators, type of data, etc) are described according to the goals of the study.
2. Life cycle inventory: Input and output flows of matter and energy as well as emissions to the environment (air, water, soil emissions and solid wastes) included in the system are listed.
3. Life cycle impact assessment: Inventory flows are converted into potential environmental impact categories using a characterization method. Each flow can contribute to several environmental impact categories. Impact categories and associated characterization method are chosen in accordance with the goals and scope of the study.
4. Interpretation of results: Results are analyzed regarding the defined goal and scope of the study.

This methodological framework is also clarified in the ILCD Handbook [21] that provides further guidance to assure consistency and quality of LCA studies.

There are two main approaches adopted in LCA studies depending on the type of questions the authors want to answer: Attributional LCA (A-LCA) and Consequential LCA (C-LCA). In an A-LCA, all the flows physically linked to the product's life cycle are included in the system's boundaries [22]. C-LCA has emerged as a modeling approach that captures impacts occurring beyond direct physical relationships assessed in A-LCA [22]. It extends the system's boundaries compared to A-LCA in order to consider market information in the life cycle inventory to assess the effects of a decision on the system [23].

LCA results could also vary from one study to another because of different sources of uncertainties. The nature of these uncertainties could be stochastic uncertainties (i.e. uncertainties linked to values of process data or characterization factors for example) or choice uncertainties (i.e. choice of methodological assumptions, impact assessment method, system boundaries, localization of data, etc) or lack of knowledge of studied system [21]. Uncertainties should be addressed in LCA studies by applying for instance Monte Carlo method to assess stochastic uncertainties or by conducting a sensitivity analysis to assess choice uncertainties.

#### 2.1.1 Specificities of LCA applied to biofuel pathways

The first applications of LCA for the environmental evaluation of biofuels were carried out in the 90's and since then; many methodological issues concerning this product category have been emphasized. The main specific methodological assumptions on biofuel LCA studies are:

- *system boundaries*: usually, a distinction is made between "Well To Tank" (WTT) boundaries that include all steps from the production of biomass feedstock to the transport and distribution of fuel and "Well To Wheel" (WTW) boundaries that include the WTT steps and the fuel use (end-of-life). Infrastructures may or may not be included within the system boundaries.
- *functional unit*: it is a measure of the function of the studied system. All LCA results from the same study should be expressed in the same functional unit to enable comparison. A usual functional unit in LCA of transportation systems is a "kilometer driven by a reference vehicle on a standard driving cycle (and assuming that generally the different fuels have a similar performance in terms of acceleration, max speed, etc.)". Another classical functional unit for assessing fuels is "the consumption of one MJ of fuel in a motor" expressed in MJ.
- *reference system*: results of the studied system have to be compared with results of a reference system (usually a fossil fuel). This reference system has to be defined in accordance with study purposes and methodological choices; in particular it must have similar boundaries, the same functional unit and similar geographical and temporal context.
- *the method to account for coproduct*: Another classical methodological issue in LCA concerns the fact that more than one product can be produced in the studied system (called coproducts). Distributing environmental burdens among products and coproducts of a process is a controversial issue in LCA. Two types of methodology are generally applied for the multiproduct cases: the substitution method and allocation method. This last method consists in sharing proportionally the environmental impacts between products and coproducts based on physical (e.g. mass, energy) or economical characteristics of the products. With the substitution method, allocation is avoided and the burdens associated to alternative ways of producing the coproduct are subtracted from the final result. The LCA ISO standards recommend the system expansion method (also called substitution method) [24,25] but the choice of the method to account for coproducts strongly depends on the purpose of the study and on the nature of the studied system.

Biofuels use biomass as raw materials. Hence, LCA applied to biofuel pathways has to deal with some classical issues linked with the biomass production:

- *Land Use Change (LUC)*: It refers to all changes induced by land conversion or land management changes. Direct LUC is mainly treated as the above and below ground carbon release from the conversion of forests or grasslands into agricultural land. Indirect LUC refers to all changes that occur when the increased demand for agricultural products induces land conversion in other parts of the world. It is important to note that these changes not only affect GHG emissions but other environmental aspects such as biodiversity, soil fertility, etc. Indirect LUC is the main subject of debate nowadays concerning biofuel environmental assessment, especially regarding GHG emissions [26] but there is no consensus on the way how to account for it in LCA methodology.
- *Nitrogen cycle*: Nitrous oxide (N<sub>2</sub>O) field emissions are known to be the subject of controversy in the biofuel LCA world since Crutzen *et al.* [27] published "N<sub>2</sub>O release from agro-biofuel production negates global warming reduction by replacing fossil fuels". There is a huge uncertainty about these emissions because they depend on local factors and this gas has a high GWP (around 300 times as much as CO<sub>2</sub>). In a G1 biofuel LCA study conducted for the French government, the uncertainty on these emissions is estimated to be 50% [28]. To estimate these emissions, some studies use the IPCC Tier 1 methodology [29] based on the amount of nitrogen fertilizer applied in the culture. However, N<sub>2</sub>O emissions depend on other factors such as soil characteristics and climate. Other assessment methods including these factors should provide a more accurate estimation.
- *Carbon cycle*: Considering the short-term carbon cycle, many biofuel LCA studies suppose that the amount of carbon captured by the biomass during the photosynthesis is equal to the amount of carbon released in the atmosphere during the biofuel combustion. So those studies do not take into account either the carbon stored by the biomass or the carbon releases during biofuel use, this is called the carbon-neutrality hypothesis.

## 2.2 General presentation of *meta-analysis* method

*"Meta-analysis refers to the statistical analysis of a large collection of results from individual studies for the purpose of integrating the findings. It connotes a rigorous alternative to the casual, narrative discussions of research studies which typify our attempts to make sense of the rapidly expanding research literature"* [30].

The Glass' pioneering articles [30–32] in educational research are usually cited in the literature as being the first ones to propose and develop this method. Over the past three decades, *MA* has first been extensively applied to clinical studies in psychological and educational research and then to health sciences. It is now increasingly employed in other research fields. Since the early 1990s, this method has been gradually more and more accepted in social sciences, such as marketing and economics<sup>3</sup>.

To our best knowledge, two articles can be identified as the precursors of the application of this methodology to synthesize LCA estimates for biofuels [33,34]. Farrell *et al.* [33] aim at estimating reliable values for the net energy and life-cycle GHG emissions of corn Ethanol in the US. They carry out a harmonization exercise on 6 studies, adjusting their methods and data to what the authors argue to be best practices. Bureau *et al.* [34], on the other hand, focuses only on the energy balance since they consider there is too much controversy involving life-cycle GHG estimations (due to uncertainties in the quantification of N<sub>2</sub>O emissions from agricultural production and indirect land use change). Rather than trying to determine best estimates, they aim at identifying the main determinants of the net energy value for G1 biofuels.

*MA* is nothing else than a particular methodology of literature reviewing. This method has not been proposed to synthesize any kind of research literature, but only studies with quantitative results: *"Meta-analysis is the analysis of empirical analyses"* [35], not theoretical ones. Applied to environmental evaluation methods, this methodology is thus relevant to review previously reported LCA studies outcomes.

Research synthesis aims at summarizing findings in such a way that clear and uncontroversial conclusions may be drawn from previous accumulated knowledge. Yet, estimates obtained by LCA approach are characterized by large differences among study results. Even if different studies deal with a same issue, each one departs from previous literature by using different data sets, different methodological choices, etc. Research synthesis may thus appear as an especially difficult task when reviewing LCA literature. Compared to qualitative literature reviews, the original idea behind *MA* is to consider study results in the same way as any scientific phenomenon. Each reported result is viewed as an "observation" of a complex dataset, *"no more comprehensible without statistical analysis than would hundreds of data points in one [LCA] study"* [32]. *MA* may then be understood as a set of statistical techniques, such as econometrics, which allows to systematically summarize quantitative studies. It is a complementary method to narrative literature surveys that generally provide a more qualitative than quantitative analysis of estimate results. Using econometrics methods, *MRA* allows to review and analyze previous results through a *ceteris paribus* reasoning [36]. By doing so, outcomes from many studies can be integrated and combined in such a way that comparison between their results become easier.

*MA* provides a quantitative summary of estimates results, such as mean estimates and confidence intervals of the quantitative results among studies. Compared to narrative literature surveys, the major contribution of *MA* consists in modelling estimate result variations as a function of different factors. The use of specific econometrics method allows then to statistically estimate and quantify their influence on study outcomes.

More formally, let the generic form of the linear regression model be the "original model" of the *MRA* equation:

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<sup>3</sup> Six meta-analyses were published at the same time in the field of economics [35,106–110]. See for instance Stanley [40] for a more comprehensive presentation. Standard references for technical aspects of meta-analysis are [38,39,111,112].

$$\begin{aligned} Y &= f(X) + \varepsilon \\ &= X\beta + \varepsilon \end{aligned} \quad (1)$$

where  $Y$  is the  $(I \times I)$  dependent variable vector composed of the  $I$  reported estimates of the phenomenon of interest in the  $MA$ . For reasons that will be developed in Section 3.1.3, the reported estimates of a  $MA$  are named "e-s" estimates. These  $I$  estimates are drawn from  $J$  studies. Note it is generally stated  $I \geq J$ . If only one estimate per study is retained, then  $J = I$ . As usual, the term  $\varepsilon$  is a  $(I \times I)$  vector of a random disturbance. It is assumed that the sampling error is normally distributed with mean zero and variance  $\sigma_{\varepsilon,i}^2 : \varepsilon_i \sim N(0, \sigma_{\varepsilon,i}^2), \forall i = 1, \dots, I$ .  $X$  is the  $(I \times K)$  matrix composed of the  $K-1$  independent variables of this meta-model. The independent variables represent study characteristics which are supposed to have an influence on the systematic excess variation of  $Y$ .  $\beta$  is the  $(K \times I)$  vector of the coefficients of this meta-model. Once estimated, it gives a measure of the particular effects of each characteristic.

The following notational convention will apply in the remaining of this paper: let the first column of the  $(I \times K)$  data matrix,  $X_{(I,K)}$ , be a column of 1s and the others column vectors be the  $I$  observations of the  $K-1$  independent variables:

$$X_{(I,K)} = \left( \begin{array}{c|c|c|c|c} C & X_1 & \dots & X_l & \dots & X_{K-1} \\ \hline (I,1) & (I,1) & & (I,1) & & (I,1) \end{array} \right) \quad (2)$$

$$\text{where } C_{(I,1)} = \begin{pmatrix} 1 \\ \vdots \\ 1 \\ \vdots \\ 1 \end{pmatrix} \text{ and } X_l_{(I,1)} = \begin{pmatrix} x_{l,1} \\ \vdots \\ x_{l,i} \\ \vdots \\ x_{l,I} \end{pmatrix}.$$

Let us specify the  $(I \times I)$  vector of the coefficients,  $\beta_{(I,1)}$ , as follow:

$$\beta_{(K,1)} = \begin{pmatrix} \alpha \\ \beta_1 \\ \vdots \\ \beta_l \\ \vdots \\ \beta_{K-1} \end{pmatrix}.$$

According to this notational convention,  $x_{l,i}$  is the  $i$ -th observation of the  $l$ -th independent variable ( $i = 1, \dots, I$  and  $l = 1, \dots, K-1$ ).  $\beta_l$  is the coefficient of the vector of the  $I$  observations of the  $l$ -th independent variable,  $X_l$ , and  $\alpha$  is the constant term in the model, also known as the intercept.

It may be convenient to refer to a single observation in eq. (1). Then, eq. (1) may be rewritten as follows:

$$\begin{aligned} y_i &= \alpha + \beta_1 x_{1,i} + \beta_2 x_{2,i} + \dots + \beta_l x_{l,i} + \dots + \beta_{K-1} x_{K-1,i} + \varepsilon_i, \quad \forall i = 1, \dots, I \\ &= \alpha + \sum_l^{K-1} \beta_l x_{l,i} + \varepsilon_i, \quad \forall i = 1, \dots, I \\ &= X_i'_{(1,K)} \beta_{(K,1)} + \varepsilon_i, \quad \forall i = 1, \dots, I \end{aligned} \quad (3)$$

where:

$$X_{(I,K)} = \begin{pmatrix} X'_1 \\ (1,K) \\ X'_2 \\ (1,K) \\ \vdots \\ X'_i \\ (1,K) \\ \vdots \\ X'_I \\ (1,K) \end{pmatrix} \text{ and } X'_i = (1, x_{1,i}, \dots, x_{l,i}, \dots, x_{K-1,i}), \forall i = 1, \dots, I$$

In *MRA* dealing with LCA studies,  $X$  could be stated as being composed of three kinds of variables.  $X = (C, T, M, S)$  where  $T, M$  and  $S$  are assumed to be  $(I \times t)$ ,  $(I \times m)$  and  $(I \times s)$  vectors, respectively.  $T$  is composed of  $t$  variables related to technical characteristics of pathways assessed in the primary studies. In this *MRA*, it corresponds to biofuel characteristics such as the type of biomass feedstock, the type of technologies and associated yields, etc. The  $m$  variables of  $M$  refers to methodological assumptions reflecting researcher choices: for instance the type of LCA approach (A-LCA or C-LCA), the system boundaries, etc. Finally, the  $s$  variables of  $S$  correspond to the typology of the study under consideration such as the type of this study (peer review or working paper for instance), the publication year or the geographical location of authors. Of course, the definitive specification of eq. (1) depends on both the particular issue investigated (here, Global Warming impact indicator of advanced biofuels) and studies reviewed in the *MRA*.

### 2.2.1 Treatment of heteroskedasticity in meta-regression analysis

Heteroskedasticity is a well-known problem in *MRA* literature. Recall that the basic linear regression model assumes  $E(\varepsilon\varepsilon') = \sigma_\varepsilon^2 I$ . This assumption implies that the variance-covariance matrix of the vector of parameters estimates,  $\beta_{(K,1)}$ , is equal to  $\sigma_\varepsilon^2 (X'X)^{-1}$ . More particularly, it is thus assumed:  $\sigma_{\varepsilon,i}^2 = \sigma_\varepsilon^2, \forall i = 1, \dots, I$ . When applied to the *MRA* framework, the homoskedasticity assumption of the disturbances may not be held.

By nature, primary studies results are not estimated with the same precision. In econometric terms, it means that each estimate has a different standard error, that is:  $\sigma_{\varepsilon,i} \neq \sigma_{\varepsilon,j}, \forall i \neq j$ . As a consequence, the variance of  $\varepsilon$  in eq. (1) varies across its observations and  $e$ -s estimates,  $y_i$ , may not be considered as having homogeneous variances. Indeed, " $e$ -s" estimates are drawn from different primary studies. These studies use different *i*) technical data/characteristics, *ii*) author's methodological choices and *iii*) do not have the same typology. These reasons, among others, may explain why each  $e$ -s estimates are estimated with varying degrees of precision.

In presence of heteroskedasticity, the Ordinary Least Square (OLS) estimates,  $\beta_{(K,1)}$ , remain unbiased and consistent. Nevertheless, heteroskedasticity often leads to wider parameter estimate confidence intervals which may cause insignificant relationships between independent and dependent variables if not accounted

for<sup>4</sup>. Therefore, heteroskedasticity is potentially a serious problem and has to be explicitly treated in *MRA*. Various solutions have been used in the *MRA* literature to correct for heteroskedasticity<sup>5</sup>. Two major approaches have been employed in the literature.

## 2.2.2 Methods of estimation using Heteroskedastic Consistent Covariance Matrix

One of the most common approaches is to use heteroskedastic consistent estimators such as White's or Huber-White's Heteroskedastic Consistent Covariance Matrix (HCCM). The Newey-West estimator has also been used in some *MA*. The latest has been designed for stationary time-series data and, as a consequence, Nelson & Kennedy [37] do not recommend to employ this estimator in a *MRA* framework. The use of White and/or Huber-White standard errors theoretically corrects for heteroskedasticity.

Nevertheless, non homogeneous variances may remain in practice, more particularly when *MRA* are applied to small sample sizes. The white and Huber-White estimators are generally used because the source of heteroskedasticity is not exactly known. It is not the case in the context of *MRA* in which the source of heteroskedasticity is clearly identified. Indeed, it has already been explained that *MRA* are subject to heteroskedasticity because e-s estimates are obtained with varying degrees of precision. That is to say, their respective standard errors are not the same. In economic sciences, e-s estimates correspond to partial regression coefficients drawn from primary studies. When estimating these coefficients, primary studies also estimate their standard errors. These estimates provide a measure of the *MRA* heteroskedasticity. This information may be used to adequately correct for heteroskedasticity. The Weighted Least-Squares (WLS) method of estimation takes such information explicitly into account in its estimation procedure.

## 2.2.3 The weighted least-squares method of estimation

A second alternative consists in estimating the parameters by using the WLS regression. Indeed, if  $y_i$ 's variances are known, the most straightforward method of the correction of heteroskedasticity is by means of WLS<sup>6</sup>.

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<sup>4</sup> A wider confidence interval of a coefficient, say  $\beta_i$ , means that its variance,  $\sigma_{\beta_i}^2$ , is greater than expected. Thus, it conducts to

a decrease of the *t-value* of  $\beta_i$ ,  $t_{\beta_i} = \frac{\hat{\beta}_i}{\sqrt{\sigma_{\beta_i}^2}}$ , which increases the probability of falsely accepting the null hypothesis of tests

of significance.

<sup>5</sup> See for instance Nelson & Kennedy [37] for a review of heteroskedasticity treatments used in meta-analysis studies dealing with environmental economics issues.

<sup>6</sup>As explained in Gujarati [113], once the original model has been transformed, the variance of "new" disturbance terms,  $\varepsilon_i^*$ , is:

$$\begin{aligned} \text{Var}(\varepsilon_i^*) = E(\varepsilon_i^{*2}) &= E\left(\left(\frac{\varepsilon}{\sigma_{\varepsilon,i}}\right)^2\right) \\ &= \frac{1}{\sigma_{\varepsilon,i}^2} E(\varepsilon) \quad \text{since } \sigma_{\varepsilon,i}^2 \text{ is known} \\ &= \frac{1}{\sigma_{\varepsilon,i}^2} (\sigma_{\varepsilon,i}^2) \quad \text{since } E(\varepsilon) = \sigma_{\varepsilon,i}^2 \\ &= 1 \end{aligned}$$

which is a constant. That is, the variance of the transformed error term,  $\varepsilon_i^*$ , is now homoskedastic.

Let  $\sigma_{\varepsilon,i}$  be the estimated standard error<sup>7</sup> of the  $i$ -th  $e$ -s estimate,  $y_i$ , for any  $i$ . Knowing the  $y_i$ 's heteroskedastic variances,  $\sigma_{\varepsilon,i}^2$ , the WLS method of estimation takes this information into account explicitly by, first, dividing eq. (3) by the standard errors of  $y_i$ ,  $\sigma_{\varepsilon,i}$ , giving:

$$\frac{y_i}{\sigma_{\varepsilon,i}} = \alpha \cdot \frac{1}{\sigma_{\varepsilon,i}} + \sum_l^{K-1} \beta_l \frac{x_{l,i}}{\sigma_{\varepsilon,i}} + \frac{\varepsilon_i}{\sigma_{\varepsilon,i}}, \quad \forall i = 1, \dots, I \quad (4)$$

Second, the Ordinary Least-Squares (OLS) method of estimation is applied to the transformed variables, *i.e.* to eq. (4).

The more  $\sigma_{\varepsilon,i}$  is important, the less is the precision of  $y_i$ . Thus, by dividing each  $y_i$  by its standard error estimate,  $\sigma_{\varepsilon,i}$ , the WLS allocates to each  $e$ -s estimate a weight which is inversely proportional to its degree of precision. Intuitively, less precise  $e$ -s estimates,  $y_i$  with wider  $\sigma_{\varepsilon,i}$ , obtain relatively smaller weight than more precise ones in minimizing the (weighted) sum of residual squares. Indeed, recall that the OLS method consists of minimizing the sum of residual squares:

$$\text{Min} \sum_{i=1}^I e_i^2 = \text{Min} \begin{pmatrix} e' & e \\ (I,1) & (I,1) \end{pmatrix}$$

where  $e_{(I,1)}$  is the column vector of residuals defined as follows:

$$\begin{aligned} Y_{(I,1)} &= X_{(I,K)} \beta_{(K,1)} + e_{(I,1)} \\ \Leftrightarrow e_{(I,1)} &= Y_{(I,1)} - X_{(I,K)} \beta_{(K,1)} \end{aligned}$$

where  $\beta_{(K,1)}$  is the column vector of parameters estimated by the OLS method.

Thus, applying the OLS method to eq. (4), WLS parameters estimates are obtained by minimizing:

$$\begin{aligned} &\text{Min} \sum_{i=1}^I \left( \frac{e_i}{\sigma_{\varepsilon,i}} \right)^2 \\ \Leftrightarrow &\text{Min} \sum_{i=1}^I \frac{1}{\sigma_{\varepsilon,i}^2} \cdot e_i^2 \quad (5) \end{aligned}$$

$$\Leftrightarrow \text{Min} \sum_{i=1}^I w_i e_i^2 \quad (6)$$

According to eq. (5) and (6), the WLS estimators are obtained by minimizing a weighted sum of residual squares with the  $y_i$ 's unconditional variances acting as the weights<sup>8</sup>:

$$w_i = \frac{1}{\text{Var}(y_i)} \quad (7)$$

Weights defined in eq. (23) are known as being those that minimize the variance of the WLS estimators. These weights will then provide estimators that are BLUE (Best Linear Unbiased Estimators). In a particular framework of *MA* (the Fixed Effects Size model), these particular weights are obtained from the estimated standard error of each  $e$ -s estimates,  $y_i$ , drawn directly from primary studies [38,39].

<sup>7</sup> Again, like  $e$ -s estimates, estimated standard errors are drawn from primary studies.

<sup>8</sup> The comparison of eq. (4) and eq. (6) may explain some confusion encountered in the literature. As highlighted by Nelson & Kennedy, some studies refer to weights based on variances and others refer to weights based on standard errors. As shown in eq. (4), standard errors weights,  $\sigma_{\varepsilon,i}$ , are used to transform the variables, but as a consequence (see eq. (5)), it is variances weights,  $\sigma_{\varepsilon,i}^2$ , which are required to minimize the weighted sum of residual squares.

## 3 Database of LCA results of GHG emissions for advanced biofuels

### 3.1 Construction and composition of the database

As mentioned before, the goal of this study is to explain the variations of LCA results for GHG emissions of advanced biofuels. Consequently, the variable of interest (so-called *effect-size* (*e-s*) or dependent variable) is the result for GHG emissions per MJ of biofuel calculated with a LCA approach. These estimates have been drawn from the study sample of this *MA*. One value for GHG emissions (i.e. the estimate) corresponds to one observation in our *MA* sample. As one study can contain several estimates, our database (i.e. our *MA* sample) can be composed of more than one observation per study ( $I \geq J$ , recall eq.(1)).

The inclusion of all estimates from a single study is a source of disagreement in the *MA* literature. Some authors believe that only one estimate should be included per study based either on the mean of the available estimates, or selected on the basis of expert judgment, while other authors advocate including all estimates as a method of boosting sample size (see Stanley [40] for a discussion on this issue). We choose to include all estimates from a single study for the following two reasons. First, the choice of a particular estimate is subjective, and when facing the same estimates, different researchers may undoubtedly make different choices. To maintain a position as neutral as possible, we considered all available explicit results in the study or which are easily inferred. Second, the core of *MA* is to summarize quantitative literature in a systematic way regardless of its quality. Hence, it would not be relevant to select studies *ex-ante* regarding their quality since this choice would be arbitrary. The *MRA* literature proposes various *ex-ante* tests (such as statistical ones) that can lead to exclude some studies *ex-post*, or at least some of their estimates, from the database/*MA* sample.

#### 3.1.1 Selection and description of studies

Before proceeding to a *MA* analysis, the database of the *MA* has to be constituted. To do so, some common procedures exist in *MA*. Stanley [40] describes three steps to conduct a *MA*. First, primary studies having estimated a common quantitative effect are identified among published and unpublished literature. This set of studies is the material of the *MA*. Second, each article results and features are coded in a database. By doing so, studies are characterized in a way that allows them to be compared. Their findings, *i.e.* their estimates, become the observed values of the dependent and independent meta-variables. The *e-s* and potential factors which are supposed to have any influence on its variations are identified and summarized in a coded form: the explanatory variables of the matrix *X*. Third, the *MRA* can be conducted to estimate the magnitude of the quantitative effect under consideration and better understand variations in the reported estimates.

This section details the selection process of studies included in this *MA*. To obtain and analyze estimates for the GHG emissions of advanced biofuels, a large bibliographical research has been carried out to collect studies using an LCA approach. We have taken a census of both published articles and "grey literature", such as unpublished papers, conference papers, official reports. The existence of published articles which present detailed literature reviews dealing with close issues than ours has already been mentioned: [8–13]. These literature reviews were the starting point of the bibliographic research. Entries of their bibliographic references have been systematically reviewed. Then, to complete this first paper selection, a web-based keyword search - e.g. "LCA", "biofuel", "second generation biofuel", "third generation biofuel", "advanced biofuel", "cellulosic ethanol", "lignocellulosic ethanol", "synthetic diesel", "syndiesel", "BTL", "microalgae", "microalgae biodiesel", etc. - has been done on relevant literature databases (Science Direct, Web of Science, SciVerse, Springer Link, etc ), web sites of major publishers of academic journals (Blackwell, Elsevier, Kluwer, Sage, Springer, Taylor Francis, and Wiley). The "grey literature" has been more particularly collected through Google and Google Scholar, Dissertation Abstracts, web sites of key academic institutions and authors and web sites of major environmental evaluation conferences.

To better insure the homogeneity of the sample, studies have to meet three selection criteria to be included in the sample of this *MA*: i) only studies with primary results were included to avoid double counting (no literature reviews)<sup>9</sup>, ii) only studies using an LCA approach were included<sup>10</sup>, iii) only LCA studies on the following liquid transportation fuels were included: lignocellulosic ethanol, FT diesel, microalgae HVO and FAME<sup>11</sup>, iv) only studies assessing Global warming impact indicator (i.e. GHG emissions) with "Well To Tank" (WTT) or "Well To Wheel" (WTW) boundaries<sup>12</sup>. The proxy used to measure the GHG emissions has to be expressed (or easily convertible) in term of grams of CO<sub>2</sub> equivalent per MJ of biofuel.

Moreover, no a priori filter was used concerning the type of publication (published or unpublished papers) but the date and the English language. This *MA* focuses on studies conducted since 2002 (until mid 2011) since, to our knowledge, no advanced biofuels LCA studies were conducted before this date.

At the end of this selection process, the database contains 47 LCA studies [4,5,41–84] providing 593 estimates of life-cycle GHG emissions of advanced biofuels. Details of number of estimates by studies included in the sample are provided in Table 2 (see Table B.1 in Appendix B for details about selected studies).

### 3.1.2 Choice and description of the meta-variables

The object of this *MA* is twofold.

First, this *MA* proposes a statistical summary of the role of different determinants for estimates of the *e-s*, i.e. the Global warming impact indicator for advanced biofuels in grams of CO<sub>2</sub>eq per MJ. By identifying and measuring the influence of these determinants, one may obtain a more in-depth explanation of how advanced biofuel LCA GHG emission estimates change as these factors vary. Second, an important aspect of this article is to provide average estimates of the Global warming impact indicator for advanced biofuels. The dependent (*e-s*) and independent variables (potential factors) of this *MA* are now detailed.

### 3.1.3 The *Effect-Size*: the dependent variable

As mentioned before, the variable of interest, so-called *effect-size* (*e-s*) or dependent variable, is the result for GHG emissions per MJ of biofuel calculated with an LCA approach. Those estimates drawn from studies, i.e. the observations of our *MA* sample, are expressed in different units of measure. These values need to be converted in a way that allows them to be combined to constitute the meta-dependent variable. The transformation of the dependent variable observations into a unique metric measure is a common procedure of *MA* studies. This step is called the *e-s* calculation and is central to *MA* literature. Indeed, it is this conversion of the dependent variable in a standard measure, the *e-s*, that allows to compare previous results and to investigate their determinants. In our sample, most of the studies present the GHG emissions, in grams of CO<sub>2</sub> equivalent, as a midpoint impact category using IPCC's characterization factors. Some other studies present only inventory data on GHG emissions so these results had to be converted into grams of CO<sub>2</sub> equivalent. We used the latest IPCC characterization factors [85] for these conversion steps. It was not possible to harmonize all of the observations by using the IPCC's 2007 characterization factors because the decomposition in individual GHG emissions were not always presented. However it has been shown that the calculation method for global warming impact has an insignificant influence in LCA results [86,87].

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<sup>9</sup> The *MA* literature distinguishes primary studies from secondary ones. Compared to the latter, the former presents original research results. Literature reviews are the typical example of secondary studies. In order to avoid double counting, only results drawn from primary studies are included in a meta-database.

<sup>10</sup> Only studies following the ISO 14044 guidelines to conduct an LCA were included [20]

<sup>11</sup> Studies on other biomass derived fuels such as methanol, DME, ETBE, biogas, heat, power, CHP were not included.

<sup>12</sup> To be more precise, only the WTW studies with consumption of pure biofuel (E10, B10, etc) have been included. No study with a bi-functional unit was included.

However, there is still another step in the calculation of the e-s since the LCA results are not always presented for the same functional unit. Typical functional units in biofuel LCA studies are a unit of fuel produced (liter, kg, MJ, etc.) or the service rendered by the biofuel (dislocation of a vehicle for a certain distance expressed in km, miles, etc.). Some other studies present their results using other less conventional functional units such as the surface of arable land used. All of these choices depend on the initial goals of the study.

We choose to convert the GHG emission values in our database into a common functional unit, a MJ of fuel produced since this is the unit used in the RED (the RFS also presents results for biofuel energy content, in Btu). For a given study, we apply conversion factors using the provided information in the study for lower heating values (LHV), densities, engine fuel consumption, etc. Whenever these values did not appear in a study, information from a well-documented study was used [88]. Some studies had to be discarded because results were presented for a functional unit that could not be converted into a MJ (e.g. Melamu *et al.* [89], is a C-LCA study where the results are presented for a multi-functional unit, involving fuel and electricity production).

Lastly, a standard error is associated to every observation in order to be able the treatment of our sample for heteroskedasticity. As mentioned before, there are mainly two ways to treat uncertainty in LCA (and consequently estimate standard errors): Monte-Carlo analysis and sensitivity analysis. The standard error could be directly inserted in the database only for the observations from studies performing Monte-Carlo analysis. We calculated a standard error from the e-s variance of each sensitivity analysis performed (one study can present the sensitivity of LCA results for variations of more than one parameter, each performed separately). For the studies that did not assess the uncertainty of their results, we calculated the standard error based on all the available observations for a same type of fuel.

### **3.1.4 The potential factors: the independent variables**

There are no guidelines concerning exactly which variables, potentially influencing LCA results, have to be included in a *MA* independent variable set. Like any other scientific investigation, this choice is determined by the available data [37], LCA practitioner knowledge (see section 2.1) and the specificities of each technology (see Appendix A). Some non-intuitive variables are also included in the database. In addition, some study characteristics (country, year of publication, etc.) were included to account for potential publication biases.

Primary studies highlight different determinants of advanced biofuel GHG emission estimates whereas surveys offer a more in-depth discussion on their likely influences. According to the introduction of this section, three categories of potential determinants of GHG emission estimates are kept: technical data, methodological choices of authors and typology of the study under consideration. The latter variables are more particularly based on typical variables employed in previous *MA*.

The three categories of explanatory variables are broken down further as follows. Each category could be divided into subcategories (see Table 1). Those subcategories could gather from 2 to 18 variables. All variables are encoded either as binary – a.k.a. dummy or qualitative – variables or as quantitative variables. At present, more than 80 variables are available in the database.

**Table 1 - List of categories and subcategories of variables included in the database**

Technical data	Methodological choices	Typology of the study
Type of biofuel	Type of LCA approach	Type of study
Type of biomass feedstock	System boundaries	Year of publication
Type of coproducts	Method for taking into account coproducts	Geographical location of authors
Type of technologies and associated yields	Carbon neutral	
Geographical location of the case study	Characterization method for impact assessment	
	Method for assessing N <sub>2</sub> O emission from N input	
	Method for taking into account Land Use Change	
	Method for taking into account uncertainties	
	Number and type of environmental impact indicator assessed in the study	

A brief description of all subcategories for all categories follows (See Table B.2 in Appendix B for a comprehensive variable description and their respective name):

**Technical data.** The type of biofuel (Biomass To Liquid, Ethanol, Fatty Acid Methyl Ester or Hydrotreated Vegetable Oil) as well as the biofuel generation (G2 biofuel for BtL and Ethanol; G3 biofuel for FAME and HVO) are set as variables.

In the "type of biomass feedstock" category, due to the variety of feedstock used for biofuel production in our sample, we created groups for biomass having similar characteristics (e.g. poplar and eucalyptus are coded as farmed wood, corn stover and wheat straw are coded as agricultural residues, etc.). An additional variable was created in order to test the difference of using cultivated resources (energy crops and farmed wood) and waste / residues as feedstock (biomass from agricultural or forestry residues) on LCA results.

In the "type of technologies and associated yields" category, all different types of processes for biomass pretreatment and for conversion into fuel that we found in the literature were set as variables for BtL and Ethanol technologies. The "Mass yield provided" variable indicates if a value for a mass yield of the biofuel process unit is available in the study (this can be seen as a quality indicator for a given study) and the "Value of mass yield" indicates this value only for G2 biofuels. For G3 biofuels, we choose the daily productivity and the oil content of microalgae as quantitative variables since they have been often identified in the literature as the most influencing factors for life cycle GHG emissions of G3 biofuels. In addition, the fact to grow microalgae in open ponds or photobioreactors is set as a variable.

**Methodological choices.** All classical methodological choices for LCA are set as variables. We differentiate LCA studies with an attributional approach from LCA studies with a consequential approach (see section 2.1)

Some hypothesis relative to system boundaries are set as variables: we distinguish WTT from WTW studies and the inclusion, or not, of infrastructures within the system boundaries is also taken into account.

As highlighted in Section 2.1, the methods used to account for coproducts can have a great influence in biofuel LCAs. Therefore they were also set as independent variables. We classify the observations as either using an allocation method (based on energetic, mass content, market value, etc.) or system expansion method. Some studies mix both methods, which we call hybrid method.

The carbon-neutrality hypothesis is very common in G1 and G2 biofuel studies. However, this hypothesis is not straightforward for studies involving microalgae since they do not always capture CO<sub>2</sub> directly from the atmosphere. CO<sub>2</sub>, from flue gas for example, is generally fed into the system. Therefore, the carbon-neutrality hypothesis is set as an independent variable for G3 biofuels.

To study the influence of the choice of a characterization method for impact assessment, we make a distinction between studies that take into account 3 GHGs (CO<sub>2</sub>, CH<sub>4</sub>, N<sub>2</sub>O) and studies that take into account more than 3 GHGs.

As also mentioned in Section 2.1, N<sub>2</sub>O emissions from the field play an important role in the GHG emissions of biofuel lifecycles. The use of IPCC's method [29] or other more complex methods for estimating these emissions are set as independent variables.

Studies that take into account direct or indirect or both Land Use Change for GHG emission calculation are also identified. The method for taking into account uncertainties is identified in each study: uncertainty analysis could be conducted by a Monte Carlo analysis or by a sensitivity analysis on specific factors

(*ceteris paribus*) or no uncertainty analysis (recall section 2.1). We also try to identify if the fact that a study assess other environmental impacts than GHG emissions could influence the GHG emission results. So the number and type of environmental impact indicators assessed in the study is controlled.

**Study typology.** Other aspects than technical data or methodological choices are included in the database. The type of study is identified: it could be peer review literature or official report or legislative text (Directive or Standard) or working paper. The year of publication as well as the geographical location of the authors is also included in the database.

## 3.2 Description of the database

This *MA* covers a large portion of studies that explicitly used LCA to evaluate environmental impacts of advanced biofuels. Finally, 47 LCA studies have been selected representing 593 observations of GHG emission results representing an average of 13 observations per study (see Table 2).

Table B.1 displays a list of the studies selected for the *MA* as well as a description of some of their characteristics.

**Table 2 - List of selected studies for the MA with a description of some of their characteristics** (\* MC=Monte Carlo analysis, SA=sensitivity analysis; \*\* PR= Peer review, OR= Official Report, Dir.= legislative text (Directive or Standard), WP= Working Paper)

Study	# of Obs.	Year	E-S (mean in gCO <sub>2eq</sub> /MJ)	Type of biofuel generation	Type of LCA approach	Uncertainty analysis? (method)*	LUC?	Type of Study (PR, OR, Dir., WP)**	Geographical location of authors
Bai et al. (2010)	2	2010	27.36	G2 (Ethanol)	A-LCA	No	No	PR	Europe
Batan et al. (2010)	14	2010	-55.43	G3	A-LCA	Yes (SA)	No	PR	North America
Campbell et al. (2010)	6	2010	-9.42	G3	A-LCA	No	No	PR	Other
Cherubini et al. (2011)	6	2011	41.07	G2 (Ethanol)	A-LCA	No	No	PR	Europe
Choudhury et al. (2002)	3	2002	25.03	G2 (Ethanol & BtL)	A-LCA	Yes (MC)	No	WP	Europe
Delucchi (2006)	4	2006	39.64	G2 (Ethanol)	A-LCA	No	Yes	WP	North America
Dussault et al. (2010)	7	2010	-19.29	G2 (Ethanol)	A-LCA	No	Yes	PR	North America
Elsayed et al. (2003)	1	2003	13.00	G2 (Ethanol)	A-LCA	No	No	WP	Europe
Fazio & Monti (2011)	15	2011	16.80	G2 (Ethanol & BtL)	A-LCA	No	No	PR	Europe
Gonzales-Garcia et al. (2009a)	8	2009	114.96	G2 (Ethanol)	A-LCA	Yes (SA)	No	PR	Europe
Gonzales-Garcia et al. (2009b)	1	2009	35.39	G2 (Ethanol)	A-LCA	No	No	PR	Europe
Gonzalez-Garcia et al. (2009c)	1	2009	-9.99	G2 (Ethanol)	A-LCA	No	No	PR	Europe
Groode et al. (2007)	4	2007	9.75	G2 (Ethanol)	A-LCA	Yes (MC)	No	WP	North America
Haase et al. (2009)	2	2009	15.53	G2 (BtL)	A-LCA	No	No	WP	Europe
Hoefnagels et al. (2010)	90	2010	12.94	G2 (Ethanol & BtL)	A-LCA	No	Yes	PR	Europe
Hsu et al. (2010)	8	2010	41.89	G2 (Ethanol & BtL)	A-LCA	Yes (MC)	No	PR	North America
JEC (2007)	6	2007	11.52	G2 (Ethanol & BtL)	A-LCA	Yes (MC)	No	OR	Europe
JEC (2011)	6	2011	11.77	G2 (Ethanol & BtL)	A-LCA	Yes (MC)	No	OR	Europe
Jungbluth et al. (2007)	9	2007	61.29	G2 (BtL)	A-LCA	Yes (SA)	No	OR	Europe
Jungbluth et al. (2008)	22	2008	47.90	G2 (BtL)	A-LCA	No	No	OR	Europe
Kaufman et al. (2010)	25	2010	24.53	G2 (Ethanol)	A&C-LCA	Yes (SA)	No	PR	North America
Koponen et al. (2009)	108	2009	43.85	G2 (Ethanol)	A-LCA	Yes (SA)	Yes	WP	Europe
Lardon et al. (2009)	4	2009	94.00	G3	A-LCA	No	No	PR	Europe
Luo et al. (2009)	9	2009	163.84	G2 (Ethanol)	A-LCA	No	No	PR	Europe
McKechnie et al. (2011)	6	2011	-55.88	G2 (Ethanol)	A-LCA	No	No	PR	North America
Mehlin et al. (2003)	2	2003	8.28	G2 (BtL)	A-LCA	Yes (SA)	No	WP	Europe
Mu et al. (2010)	19	2010	-5.33	G2 (Ethanol & BtL)	A-LCA	Yes (SA)	No	PR	North America
Mullins et al. (2010)	10	2010	41.10	G2 (Ethanol)	A-LCA	Yes (MC)	Yes	PR	North America
RED (2009)	10	2009	12.80	G2 (Ethanol & BtL)	A-LCA	No	No	OR/Dir.	Europe
RFS2 (2010)	12	2010	20.67	G2 & G3	C-LCA	Yes (MC)	Yes	Dir.	North America
Sander et al. (2010)	1	2010	-18.40	G3	A-LCA	No	No	PR	North America
Schmitt et al. (2011)	3	2011	49.62	G2 (Ethanol)	A-LCA	No	No	PR	North America
Sheehan et al. (2004)	1	2004	-81.28	G2 (Ethanol)	A-LCA	No	Yes	PR	North America
Spatari et al. (2005)	2	2005	18.94	G2 (Ethanol)	A-LCA	No	Yes	PR	North America
Spatari et al. (2009)	34	2009	-2.69	G2 (Ethanol)	A-LCA	Yes (MC & SA)	Yes	PR	North America
Spatari et al. (2010)	6	2010	-7.93	G2 (Ethanol)	A-LCA	Yes (MC)	Yes	PR	North America
Stephenson et al. (2010a)	17	2010	12.12	G2 (Ethanol)	A-LCA	Yes (SA)	No	PR	Europe
Stephenson et al. (2010b)	31	2010	201.15	G3	A-LCA	Yes (SA)	No	PR	Europe
Stichnothe et al. (2009)	18	2009	33.98	G2 (BtL)	A-LCA	Yes (SA)	No	PR	Europe
Stratton et al. (2010)	23	2010	24.60	G2 & G3	A-LCA	Yes (SA)	Yes	WP	North America
van Vliet et al. (2009)	5	2009	-15.78	G2 (BtL)	A-LCA	No	No	PR	Europe
Vera-Morales et al. (2009)	4	2009	55.75	G3	A-LCA	No	No	WP	Europe
Wang et al. (2010)	3	2010	13.79	G2 (Ethanol)	A-LCA	No	Yes	PR	North America
Wang et al. (2011)	3	2011	8.00	G2 (Ethanol)	A-LCA	No	No	PR	North America
Whittaker et al. (2011)	15	2011	57.50	G2 (Ethanol)	A-LCA	No	Yes	PR	Europe
Wu et al. (2005)	5	2005	14.72	G2 (Ethanol & BtL)	A-LCA	No	No	OR	North America
Xie et al. (2011)	2	2011	-59.24	G2 (BtL)	A-LCA	Yes (MC)	No	PR	North America
<b>Number of studies</b>	<b>47</b>								
<b>Number of observations</b>	<b>593</b>								
<b>Mean and repartition (weighted by observations)</b>		<b>2009</b>	<b>34.45</b>	G2 (87%) of which BtL (26%) and ethanol (61%), G3 (13%)	A-LCA (97%), C-LCA (3%)	MC (10%), SA (38%), no uncertainty analysis (52%)	LUC (51%), no LUC (49%)	PR (65%), OR (12%), Dir. (4%), WP (19%)	North America (45%), Europe (53%), Other (2%)
<b>Mean and repartition (weighted by studies)</b>	<b>13</b>	<b>2009</b>	<b>23.07</b>	G2 (87%) of which BtL (38%) and ethanol (70%), G3 (17%)	A-LCA (98%), C-LCA (4%)	MC (21%), SA (26%), no uncertainty analysis (53%)	LUC (28%), no LUC (72%)	PR (65%), OR (12%), Dir. (4%), WP (19%)	North America (45%), Europe (53%), Other (2%)
<b>Median (weighted by studies)</b>	<b>6</b>	<b>2010</b>	<b>15.53</b>						

As displayed in Table 2, the database contains 87% of studies assessing G2 biofuels (38% of studies assessing BtL and 70% Ethanol) and 17% of studies assessing G3 biofuels. Thus, among 593 observations included in the database, observations for G3 biofuels represent 13%. Other observations correspond to G2 biofuels of which 30% are BtL and 70% are Ethanol. Most of studies are based on attributional LCA approach; only 3% of observations are calculated with a consequential LCA approach. Half of the studies do not perform an uncertainty analysis on their results. Among studies that include an uncertainty analysis, 44% are performing a Monte Carlo analysis. Only 28% of studies included in the database take into account LUC (and only 4% are addressing Indirect LUC issues), representing 51% of the observations. Observations

extracted from peer review literature represent 61% of observations (65% of studies), from the official reports 9% (12% of studies), from regulatory texts 3% (4% of studies), and from working paper 25% (19% of studies).

Furthermore, the number of studies assessing GHG emissions of advanced biofuels started to strongly grow since 2007 (see Figure 2). This phenomenon could be linked with the publication of legislative texts in the EU and the US regarding mandatory GHG emission savings threshold for biofuels (respectively RED in 2009 and RFS2 in 2010).

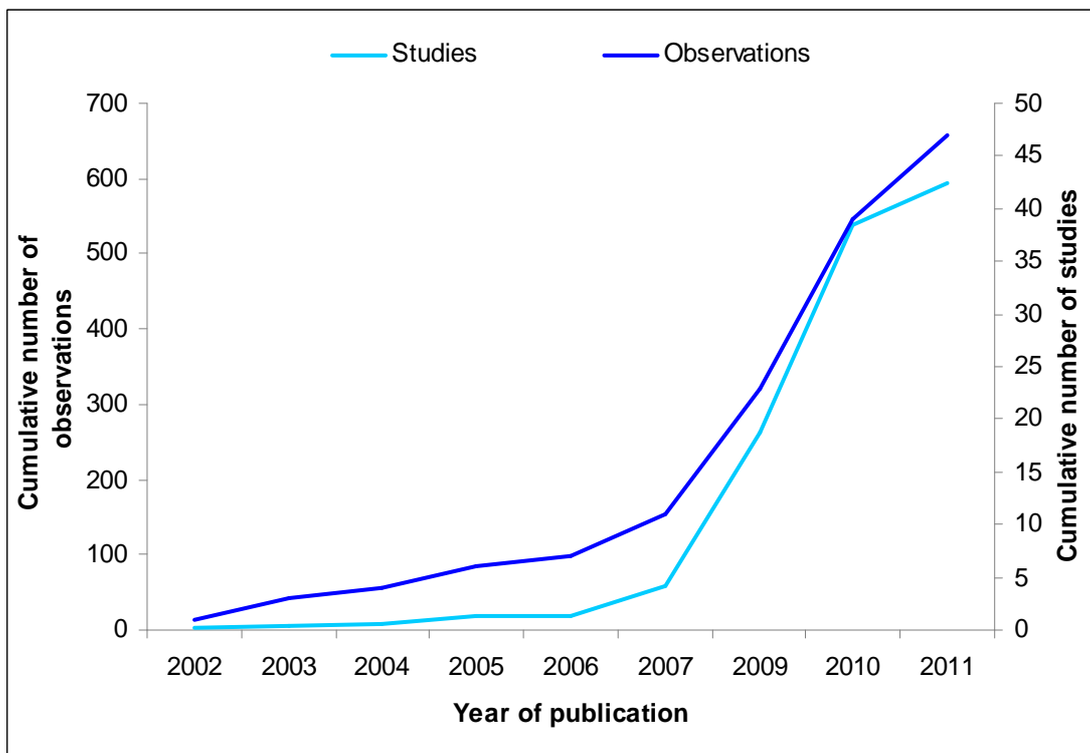


Figure 2 - Cumulative number of studies and observations per year of publication

### 3.2.1 Observations per type of biofuels

As depicted in Figure 3 (see also Table 3), GHG emission mean for G3 biofuels is quite similar to GHG emissions for fossil fuel reference as defined in EU and US regulations (respectively 83.8 g CO<sub>2</sub>eq/MJ (same reference for gasoline and diesel) and 92.5 g CO<sub>2</sub>eq/MJ (mean of US gasoline and diesel references)). GHG emission mean value indicates that G2 biofuels could induce a GHG emission reduction compared to fossil fuel reference from 69% to 72% (depending on the fossil fuel reference chosen). So, from a statistical point of view, G3 biofuels seem to emit more GHG emissions during their life cycle than G2 biofuels. In the same way, GHG emission mean for BtL is lower than for Ethanol (GHG emission savings compared to fossil fuel reference from 77% to 79% for BtL and from 65% to 68% for Ethanol).

The range of GHG emission results for G3 biofuels is very wide compared to the one for G2 biofuel as illustrated by their standard deviations (see Table 3). Hence G3 biofuel could emit 20 times more GHG emissions than fossil fuel reference whereas G2 biofuel could emit from 4 to 9 times more by considering the maximum variation of the results. Conversely, all minimum results are negative and quite similar for G2 and G3 biofuels.

Even if LCA results are inconclusive regarding GHG emission performances of advanced biofuels because of a wide range of variation, some trends can be identified: on average, GHG emissions for G3 biofuels are

higher than for G2 biofuels and GHG emissions for Ethanol are higher than for BtL. Thus, the type of biofuel seems to be a variable that could explain the differences of GHG emission results for advanced biofuels.

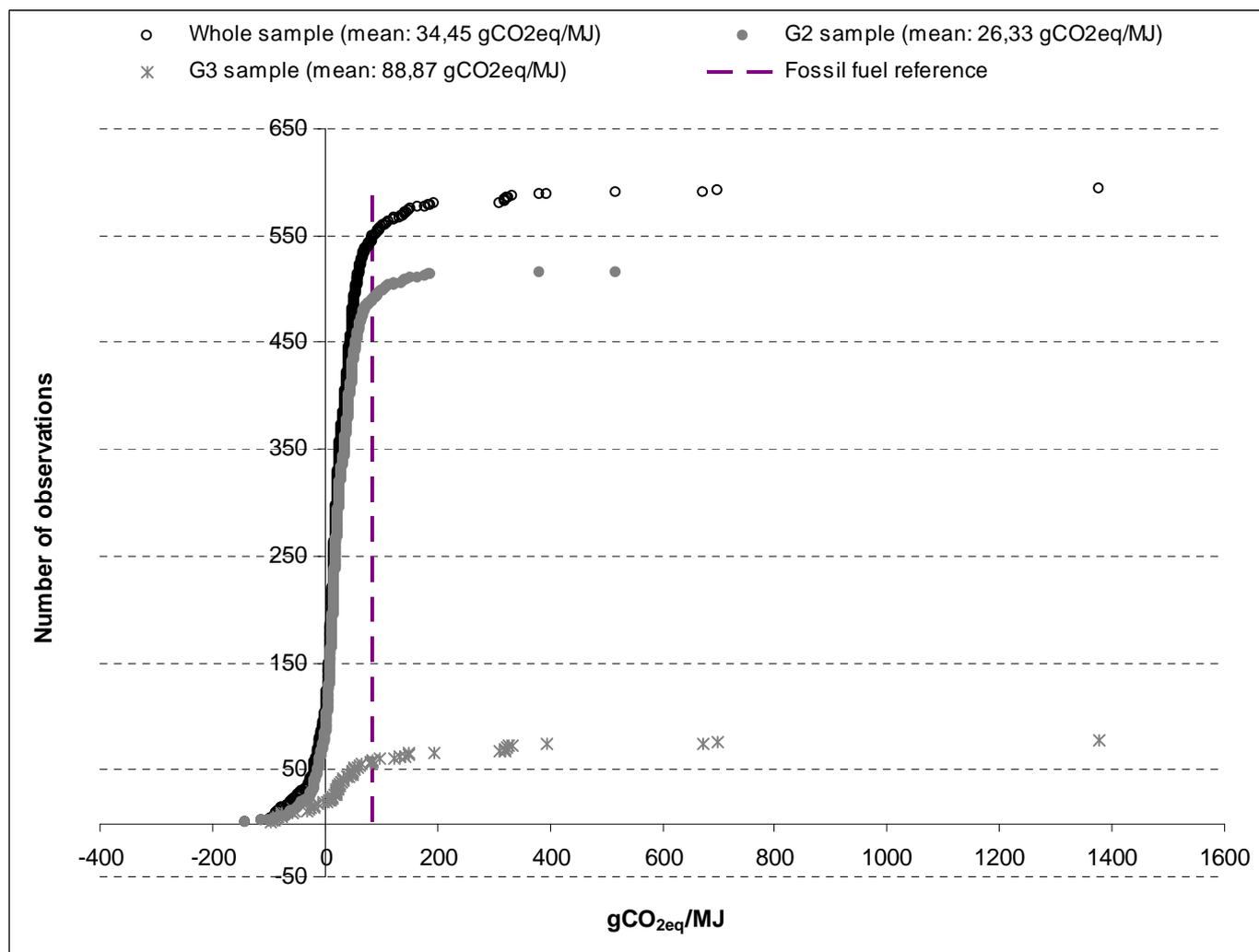


Figure 3 – Dispersion of LCA GHG emission results included in the database for the different types of biofuel

### 3.2.2 Observations per regions

We make the distinction between the geographical location of the authors (affiliation of the first author) and the geographical location of the cases studies (i.e. geographical location of inventory data).

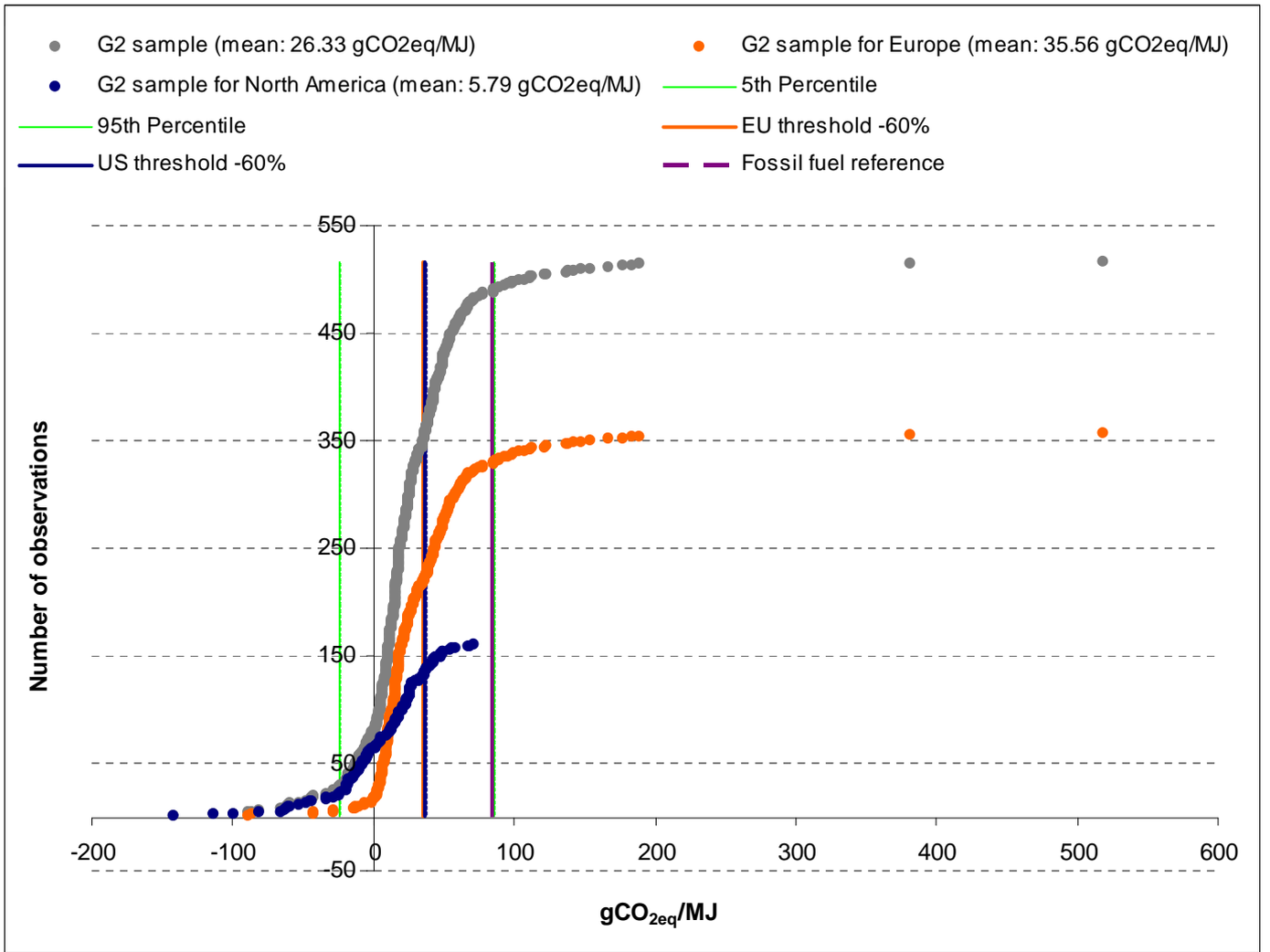
Regarding the geographical location of the authors, 45% of studies are from North American (NA) authors (including US and Canada) and 53% are from European authors (including EU countries and Switzerland), representing 32% and 67% of the observations respectively (see Table B.3 in Appendix B). The other study is from Australian authors [43]. For G3 biofuels, 42% of observations are from NA authors, 51% from European authors and 7% from Australian authors. For BtL, 23% of observations are from NA authors and 77% from European authors. For Ethanol, 34% of observations are from NA authors and 66% from European authors. In most of the studies, the geographical location of the authors fits with the geographical location of the assessed pathways. Only 3% of the observations do not match ([63] and some observations of [54]). Therefore, we focus only on the geographical location of the authors as a measure of the potential influence of geographical location on GHG emissions.

On average for all types of biofuel, GHG emission results from NA authors seem to be lower than from European authors with a gap that could be significant as illustrated in Table 3 (e.g. from 0.22 g CO<sub>2eq</sub>/MJ

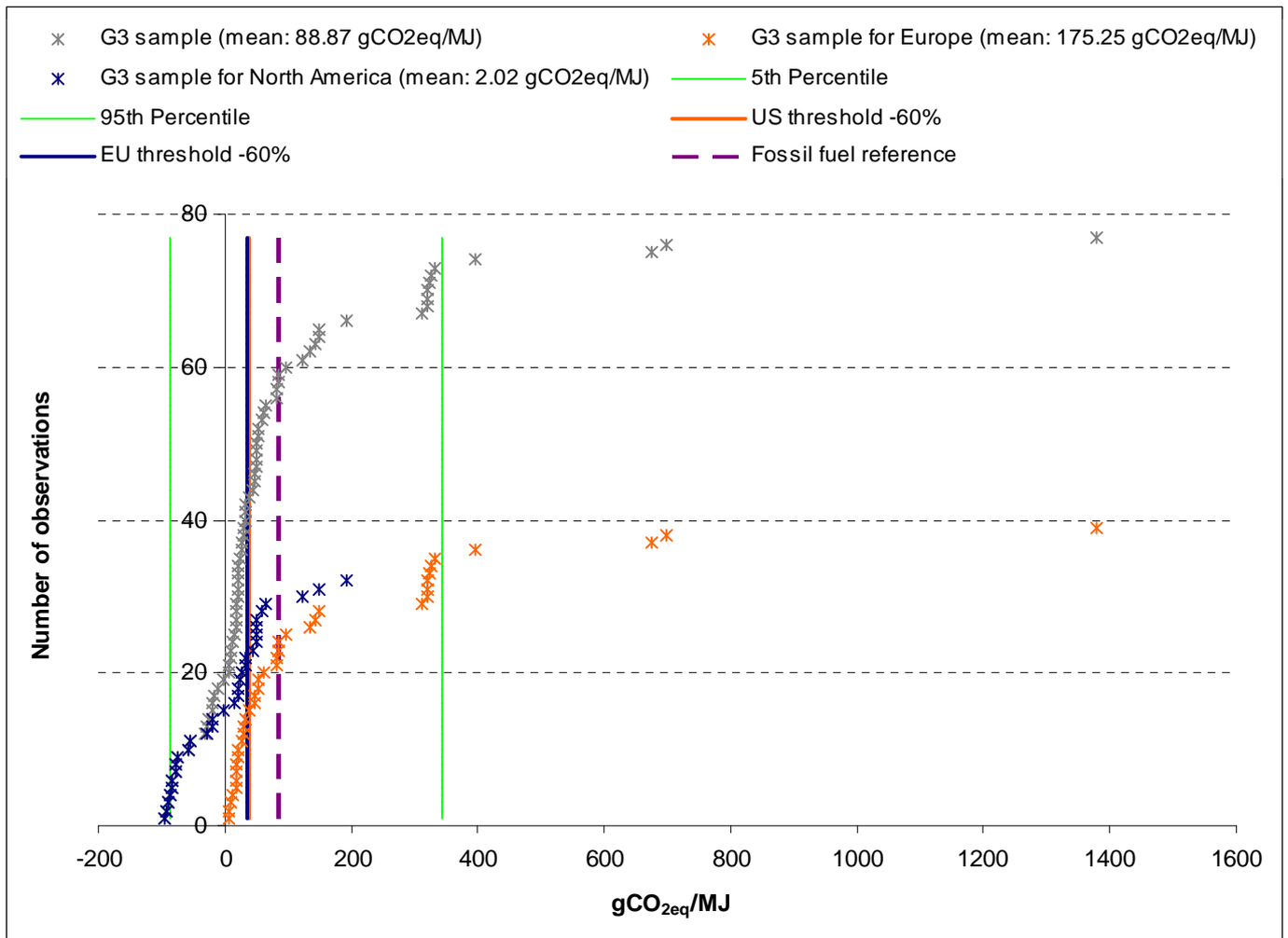
for NA to 150.63 g CO<sub>2</sub>eq/MJ for Europe for G3 biofuels). Hence, it seems that the geographical location of the authors can have an influence on the GHG emission variability observed for advanced biofuels.

**Table 3 – Statistical description of GHG emission results included in the database for the different types of biofuel and for the different geographical location of authors (\* expressed in g CO<sub>2</sub>eq/MJ)**

Biofuel generation	Location of authors	# of Obs.	(%)	Median*	Mean*	Standard deviation	Extrema*		Percentiles*	
					[Confidence Interval]		Min	Max	5th	95th
<b>G3 &amp; G2</b>	<b>All</b>	<b>593</b>		<b>21.60</b>	<b>34.45</b> [27.26;41.64]	<b>89.34</b>	<b>-142.18</b>	<b>1377.90</b>	<b>-37.08</b>	<b>116.65</b>
	North America	198	(33%)	12.61	<b>4.72</b> [-1.24;10.68]	42.78	-142.18	193.20	-79.66	55.40
	Europe	401	(68%)	26.05	<b>48.47</b> [38.53;58.41]	101.54	-88.36	1377.90	2.44	144.68
<b>G3</b>	<b>All</b>	<b>77</b>		<b>31.00</b>	<b>88.87</b> [41.55;136.19]	<b>211.85</b>	<b>-96.47</b>	<b>1377.90</b>	<b>-85.00</b>	<b>332.20</b>
	North America	38	(49%)	17.99	<b>0.22</b> [-21.62;22.05]	68.67	-96.47	193.20	-89.89	134.98
	Europe	45	(58%)	61.86	<b>150.63</b> [76.58;224.68]	253.44	-30.97	1377.90	8.69	676.39
<b>G2</b>	<b>All</b>	<b>516</b>		<b>20.50</b>	<b>26.33</b> [22.43;30.23]	<b>45.20</b>	<b>-142.18</b>	<b>518.40</b>	<b>-24.00</b>	<b>85.80</b>
	North America	160	(31%)	12.41	<b>5.79</b> [0.51;11.08]	34.12	-142.18	71.00	-60.07	49.47
	Europe	356	(69%)	24.25	<b>35.56</b> [30.72;40.39]	46.55	-88.36	518.40	1.00	100.76
<b>G2-BtL</b>	<b>All</b>	<b>155</b>		<b>14.50</b>	<b>19.04</b> [13.41;24.68]	<b>35.78</b>	<b>-142.18</b>	<b>189.00</b>	<b>-18.50</b>	<b>69.05</b>
	North America	36	(23%)	6.10	<b>-1.55</b> [-12.67;9.57]	34.05	-142.18	47.61	-54.08	32.15
	Europe	119	(77%)	15.80	<b>25.28</b> [19.16;31.39]	34.03	-88.36	189.00	2.11	85.76
<b>G2-Ethanol</b>	<b>All</b>	<b>361</b>		<b>24.30</b>	<b>29.45</b> [24.46;34.45]	<b>48.39</b>	<b>-113.60</b>	<b>518.40</b>	<b>-25.56</b>	<b>89.78</b>
	North America	124	(34%)	15.39	<b>7.93</b> [1.95;13.91]	33.97	-113.60	71.00	-61.12	49.99
	Europe	237	(66%)	30.87	<b>40.72</b> [34.22;47.21]	50.99	-42.00	518.40	1.00	104.55



**Figure 4 - Dispersion of LCA GHG emission results included in the database for G2 biofuels and for the different geographical location**



**Figure 5 – Dispersion of LCA GHG emission results included in the database for G3 biofuels and for the different geographical location**

Figure 4 and Figure 5 presents the dispersion of GHG emission results included in the database for the different types of biofuel and for the different geographical locations. These results are also compared with their respective GHG emission minimum threshold depending on their geographical location.

As already mentioned, the RED and RFS2 set minimum GHG emission savings for biofuels. Their more restrictive savings are set to 60% compared to their corresponding fossil fuel reference (fossil fuel reference are slightly different). According to Figure 3, 82% of GHG emission results from NA are compliant with their more restrictive GHG emission minimum threshold whereas only 59% from Europe are compliant with their corresponding threshold. At this stage of the analysis, we do not have objective reasons explaining this systematic difference between NA and EU estimates. It may come from the use of a different set of technical variables, for instance, but it may also reveal the existence of a potential publication bias in the literature.

In conclusion, this section based on descriptive statistics allows the formulation of some intuitions about factors that could influence GHG emission results for advanced biofuels. The type of biofuels (G2 vs G3 biofuel, BtL vs Ethanol) and the geographical location (North America vs. Europe) seem to have an influence on the variability of GHG emission results for advanced biofuels. However it is not possible to be more conclusive and accurate with the descriptive statistics presented in this Section. Descriptive statistics and inspection of graphics are very useful and often relevant but remain always vulnerable to subjective interpretation. Thus, more objective statistical tests are needed, as those that could be done with *MRA*. By using specific econometrics methods, we believe that a *MRA* should allow the *i*) confirmation of our intuitions previously identified and *ii*) to go further in the explanation of the variability by identifying and quantifying the main variation factors.

Let us now develop the *MRA* based on these LCA studies.

## 4 Results

Compared to narrative literature reviews, the *MRA* methodology allows us *i*) to statistically identify main drivers of the *e-s* variability and *ii*) to estimate both the direction and the magnitude of their respective effects across primary studies under consideration. The logic of *MRA* is illustrated here by applying this methodology to LCA literature evaluating GHG emissions of advanced biofuels. We first present the *MRA* model and its results for various G2 and G3 biofuel sub-samples. Second, we use the technique of *benefits transfer using meta-regression models* to propose a first attempt of harmonization of these LCA results.

### 4.1 The meta-regression model

Simply stated, to review a specific environmental evaluation literature, one must summarize its previous results already published on the issue under consideration.

We consider  $I$  advanced biofuels GHG emission estimates, the *e-s*, indexed by  $i = (1, \dots, I)$  and assume that the "true" *e-s* value for a given estimate is given by<sup>13</sup>:

$$y_i = \alpha + X_i' \beta + \mu_i, \quad \forall i = 1, \dots, I \quad (8)$$

where  $y_i$  is the true *e-s*,  $\alpha$  is a common factor,  $X_i'$  is a vector that measures characteristics of the biofuel case study and of the study under consideration,  $\beta$  is a vector of parameters to be estimated, and  $\mu_i$  is normally distributed with mean zero and variance  $\tau_{\mu,i}^2 : \mu_i \sim N(0, \tau_{\mu,i}^2)$

The "true" *e-s* value,  $y_i$ , is not observed. Instead, each study provides an estimated *e-s*,  $\hat{y}_i$ , so that:

$$\hat{y}_i = y_i + \varepsilon_i = \alpha + X_i' \beta + \mu_i + \varepsilon_i, \quad \forall i = 1, \dots, I \quad (9)$$

where  $\varepsilon_i$  is an error term that is normally distributed with mean zero and variance  $\sigma_{\varepsilon,i}^2 : \varepsilon_i \sim N(0, \sigma_{\varepsilon,i}^2), \quad \forall i = 1, \dots, I$

Thus we allow the "true" *e-s* and the precision of the estimated *e-s*,  $\sigma_{\varepsilon,i}^2$  to vary across estimates. The term  $\sigma_{\varepsilon,i}^2$  is known as the within-variance and varies from study to study. As already mentioned, it is usually taken as given and derived from the original estimate.

Any remaining heterogeneity between estimates is either explainable by the observable differences modeled through the moderator variables contained in  $X_i'$  or is random and normally distributed with mean zero and variance  $\tau_{\mu,i}^2$ , the between-variance.

If  $\tau_{\mu,i}^2 = 0$ , the model is referred to the fixed-effects model, and it is assumed that all heterogeneity in the "true" *e-s* can be explained by differences in study characteristics. If the between-variance is not equal to zero, the model is a REM, which is usually referred to a "mixed-effects" model because it contains observable "fixed" characteristics in  $X_i'$  as well as a random unobservable component with mean zero and variance  $\tau_{\mu,i}^2$ . The unknown variance can be estimated by an iterative (restricted) maximum likelihood process or, alternatively, using the empirical Bayes method, or a non-iterative moment estimator.

Note that the meaning of the adjectives "fixed" and "random" in the *MA* literature is different from the usual interpretation for panel data models in standard econometrics, because they refer to assumptions about the

<sup>13</sup> The following presentation is partly inspired from Ready [114].

underlying population  $e$ -s [38]. In standard econometric terms, the fixed-effects meta-estimator is equivalent to the weighted least squares (WLS) estimator using the estimated variances (derived in the primary studies) as weights and re-scaling the standard errors of the meta-regression by means of the square root of the residual variance. The random effects estimator is akin to a random coefficient model in which the within- and between-study variances are used as weights [90]<sup>14</sup>.

## 4.2 Meta-regression analysis results

Since the studies in the primary literature may use different data sets and different ways of modeling, we have good reasons to suspect the heteroskedasticity.

A common approach is to use White's Heteroskedastic-Consistent Covariance Matrix (HCCM). This estimator simultaneously corrects for heteroskedasticity and cluster autocorrelation, and hence accounts for the multiple data setup by allowing different variances and non-zero covariances for clusters of measurements from the same study. However, the White estimator is arguably rather restrictive assuming that all differences across observations and studies are observable and can entirely explain the empirical heterogeneity. In addition, the White estimator does not fully exploit all available information because it estimates the variance rather than taking it as given or recoverable from the primary studies.

The latter can be remedied by using the fixed-effects meta-estimator that we already presented. As explained above,  $\sigma_{\varepsilon,i}^2$  is a sample estimate of the standard deviation of the meta-regression errors. When this kind of measure of the heteroskedasticity is available, then Weighted Least Squares (WLS) becomes the obvious method to obtain efficient estimates of eq. (9).

We start out by presenting the results obtained for the "*whole*" sample, which includes all the G2 and G3 biofuel studies included in the meta-database. Recall that our meta-database includes variables representing *i*) technical data/characteristics, *ii*) author's methodological choices and *iii*) typology of the study under consideration. As technical data are specific to each type of biofuel, it is not possible to include this set of variables in the "*whole*" sample in order to test and quantify their respective influence. In order to capture characteristics of each biofuel generation and the type of fuel analyzed, one needs to break the "*whole*" sample into these respective sub-samples. The subsequent sections present then results for smaller samples named as follows: "*G3*", "*G2*" samples and then "*G2-BtL*" and "*G2-Ethanol*" sub-samples. Hence, the "*whole*" sample corresponds to the merge of our "*G3*" and "*G2*" samples. Note that the "*G3*" and "*G2*" samples have been cut to 90% in order to exclude outliers which may have spurious influence on econometric estimates, as it is usually done in applied econometrics. So-defined, the "*G2*" sample contains 464 observations (321 for Ethanol and 143 for BtL) and the "*G3*" sample contains 69 observations. (See Figure 4 and Figure 5 for a visual representation of "*G2*" and "*G3*" samples outliers). "*G2-BtL*" and "*G2-Ethanol*" sub-samples are a subset of the "*G2*" sample.

Results of eq. (9) are presented in Table 4 for the "*whole*" and "*G2*" samples, and Table 5, Table 6 and Table 7 provide results for the "*G2-Ethanol*", "*G2-BtL*" and "*G3*" sub-samples respectively. For each model, results are systematically reported for two different corrections for heteroskedasticity: the first estimator uses the White's Heteroskedastic-Consistent Covariance Matrix (HCCM) (as denoted by the number 1 in columns) and the second one uses Weighted Least Squares (WLS) using inverse standard error weights (as denoted by the number 2 in columns).

Unless it is indicated, all regression results are presented in reduced form. These models were chosen by the *general to specific* approach to econometrics modeling. As usual, "\*\*\*\*", "\*\*\*" and "\*\*" respectively indicate 1%, 5% and 10% significance levels and standard errors of the coefficient estimates are reported into brackets. In each column, "-" means that the variable under consideration has been first included but finally removed from the reduced form because its coefficient estimate was not statistically significant at the 10% significance levels. Regarding model information,  $N$  and  $Mean\ dep. Var$  indicate respectively the number of

<sup>14</sup> Thompson & Sharp [115] provide an overview of various estimators that allow for random-effects variation.

observations used to perform each regression and the corresponding mean of the dependent variable, i.e. the mean  $e$ -s expressed in g CO<sub>2</sub>eq/MJ of biofuel.

In all tables, the quality of regressions is checked through the following diagnostic tests. Given that the simple R-squared statistic is sensitive to the number of variables included, only the adjusted R-squared is reported (*Adj. R-squ.*). The overall fit of the regression model is assessed by the logarithm of the Likelihood (*Log-Likelihood*) and the standard Fisher test which test for joint significance. The statistic of the latter test (*F-stat.*) and the corresponding measure of its statistical probability (*P.value*) are systematically reported. The null hypothesis of this test is all coefficients but the constant one is equal to zero. Two additional diagnostic tests for the quality of the regressions (and their *P. values*) are also reported: the Skewness's asymmetric test (*Skewness*) and the Kurtosis's normality test (*Kurtosis*) of residuals. They respectively correspond to a test of skewness and nonnormal kurtosis compared with the null hypothesis of symmetry (the skewness coefficient is zero for symmetrically distributed data) and kurtosis coefficient of 3. The normality tests examine the normality of the residuals. Nonnormal residuals invalidate hypothesis tests on individual variables as these tests assume their normality, and is therefore an important consideration. All Tables also report the following two information criteria: the Akaike's Information Criterion (*AIC*) and the Schwarz's Bayesian Information Criterion (*BIC*). These two standard measures are used to allow (nonnested) model comparisons. Smaller *AIC* and *BIC* are preferred, because higher *Log-Likelihood* is preferred. Finally, in order to test and hence statistically confirm the importance of including technical data/characteristics in our models, it has been chosen to perform a likelihood-ratio test. The statistic of this test (*LR test*) and its corresponding *P. value* are reported in Table 5, Table 6 and Table 7. The line *Nested model* indicates against which model the investigated model is tested. In econometric terms, the nested model is the restricted model and corresponds to the reduced model without any technical data/characteristics.

We turn now to the comments of the results obtained for each samples and sub-samples. We only focus on the signs and significance of the estimated coefficients since the absolute magnitudes of those coefficients are not important.

#### 4.2.1 Results for the whole sample

Estimates results for the "whole" sample are presented in Table 4, columns (1aAll) and (2aAll). Eq. (9) is estimated using both the White's HCCM (column (1aAll), Table 4) and the WLS (column (2aAll), Table 4) estimators. Contrary to economic primary studies, variances are usually not reported for each estimate in LCA primary studies and have to be retrieved (recall Section 3.1.3). For each observation of the *MRA*, variances have been directly inserted in the database or calculated depending whether the observations were coming from primary studies performing Monte-Carlo analysis or sensitivity analysis, respectively. As a consequence, the database does not provide a single measure of the variance for each observation. For this reason we prefer to comment coefficient estimates obtained by OLS estimator with a White procedure – *OLS (White's HCCM)*, as indicated in last line, Table 4, – rather than *WLS*. However, we let *WLS* estimates for robustness check since they yield to similar results. The same choice is applied to the remainder of the paper to simplify the exposition.

Thus, we only comment results presented in column (1aAll), Table 4. 533 observations are included in this regression. As already explained in the previous Section, this regression only aims at testing the influence of *i*) the type of biofuels (*gen\_3*, *etha* and *btl* variables) and *ii*) the geographical location (*zlab\_us*, *zlab\_eu* and *zlab\_other*) on the  $e$ -s in order to confirm or deny what have been highlighted with the visual inspections presented in Sections 3.2.1 and 3.2.2. This may explain the rather low level of the adjusted R-squared (about 16%). As judged by the *F-stat. P. value*, the joint significance of results is accepted at the 1% significance level.

As a first comment, the econometric results displayed in Table 4 tend to confirm intuitions presented in Section 3.2, which were based on a simple visual inspection. *etha* and *btl* variables are indeed statistically significant at the 1% level and their coefficients are negative. According to these parameter estimates, GHG emissions are statistically lower for Ethanol and BtL (G2 biofuels) than for G3 biofuels (*gen\_3*) by respectively about 41 and 52 g CO<sub>2</sub>eq/MJ. These results also confirm that life cycle GHG emission

performance is better for BtL than for Ethanol. One can effectively not merge the *etha* and *btl* variables, as indicated by the Wald Test: we effectively reject the null hypothesis of this test,  $H_0$ , because  $P. Value < 0.01$  and conclude that the coefficient of *etha* is statistically different from the one of *btl*. Hence the biofuel generation is a key variable to explain the variability of advanced biofuels LCA results.

Regarding the geographical location, *zlab\_us* and *zlab\_other* variables have a negative impact on GHG emissions – their coefficients are significant at the 1% level. According to these results, GHG emissions are statistically lower when studies are from NA or from other countries (excluding NA and Europe) compared to those from Europe. Hence, the geographical location appears to have an influence on GHG emission results for advanced biofuels. There is no intuitive reason to explain the geographical influence highlighted by our results. At this step of the analysis, this result could be explained by either a model misspecification or the existence of a publication bias. The former could correspond to missing variables in our database, hence the geographical location could be a shadow variable hiding a *real* determinant. For instance, the geographical location variable could hide a set of technical data specific to one location. Unfortunately, it is not possible to include such variables in the "whole" sample model. To test this hypothesis, the "whole" sample is thus divided into G3 biofuel sample and G2 biofuel sample in order to assess specific characteristics (including technical data) of each biofuel generation.

**Table 4 – Results of MRA for the econometric samples Whole and G2 biofuels**

Samples Model	Whole 1aAll	Whole 2aAll	G2 1aG2	G2 2aG2	G2 1bG2	G2 2bG2
Constant	76.27*** (13.64)	271.74*** (23.66)	20.32*** (3.43)	27.24*** (4.72)	21.14*** (3.61)	28.43*** (5.04)
<b>Technical data</b>						
gen_3 (ref for Whole)						
etha	-41.39*** (13.14)	-220.92*** (23.77)	5.84*** (1.91)	-	5.83*** (1.81)	-
btl (ref for G2)	-52.12*** (13.36)	-215.57*** (22.59)				
mat_cult					-9.47*** (2.21)	-11.56*** (3.02)
mat_cultxdluc			-7.94*** (2.46)	-13.67*** (3.23)		
<b>Methodological choices</b>						
lca_att (ref)						
lca_cons			-33.66*** (4.79)	-40.41*** (8.63)	-34.04*** (4.9)	-39.09*** (8.37)
copval_alloc			8.96*** (1.91)	8.82** (3.62)	8*** (1.94)	6.99* (3.84)
copval_systexp (ref)						
copval_hyb			5.25** (2.38)	-	5.41** (2.69)	-
luc_dir						
luc_indir			29.97*** (6.32)	39.78*** (7.27)	29.62*** (6.34)	36.54*** (7.2)
uncer_MC			8.03** (3.45)	16.68*** (4.61)	8.04** (3.41)	17.25*** (4.58)
uncer_SA			7.78*** (2.4)	7.08* (3.63)	7.32*** (2.39)	6.69** (3.39)
uncer_ref (ref)						
impcat_nev			9.26*** (2.99)	-	7.71*** (2.71)	-
impcat_nrc			-15.01*** (2.36)	-7.31** (3.41)	-12.65*** (2.54)	-
impcat_other			-	-	0.84* (0.49)	-
impcat_gwponly (ref)						
<b>Typology of the study</b>						
zlab_us	-24.6*** (3.97)	-190.58*** (25.05)	-8.32*** (2.1)	-18.58*** (3.66)	-8.66*** (2.12)	-19.73*** (3.2)
zlab_eu (ref)						
zlab_other	-85.69*** (15.6)	-281.16*** (24.85)				
<b>Model information</b>						
N	533	533	464	464	464	464
Mean dep. Var.	28.64	17.62	24.15	25.04	24.15	25.01
Adj. R-squ.	16.30%	68.76%	37.26%	30.95%	38.33%	30.94%
Log-Likelihood	-2727.20	-3068.04	-1976.89	-2044.50	-1972.36	-2044.03
F-stat.	18.93	32.82				
(P. value)	(0,0000)	(0,0000)				
Skewness	61.27		24.57		23.56	
(P. value)	(0,0000)		(0,017)		(0,0354)	
Kurtosis	8.75		1.6		3.06	
(P. value)	(0,0031)		(0,2062)		(0,0801)	
AIC	5464.39	6146.08	3977.78	4113.00	3970.72	4114.05
BIC	5485.79	6167.47	4027.45	4162.68	4024.54	4167.87
Wald Test (P. value) for etha=btl	26.29 (0,0000)	0.22 (0,6409)				
Procedure	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS

## 4.2.2 Results for the G2 sample

Estimates results for the "G2" sample are presented in Table 4, columns (1aG2) to (2bG2). Our comments are based on results presented in column (1aG2). The adjusted R-squared is now equal to about 37%.

### Technical variables

*etha* variable is statistically significant at the 1% level and impacts positively GHG emissions for G2 biofuels. Thus GHG emissions are higher by about 6 g CO<sub>2</sub>eq/MJ for Ethanol than for BtL. The type of fuel conversion technology can thus explain the variability of GHG emission results for G2 biofuels. G2 sample is then split into "G2-Ethanol" sample and "G2-BtL" samples in order to take into account specificities of each fuel (see Sections 4.2.3 and 4.2.4, respectively).

Regarding the influence of *mat\_cult*, this variable was tested first and had a negative effect on GHG emissions for G2 (results reported in columns (1bG2) and (2bG2), Table 4). Most LCA studies do not account for upstream burdens related to residue production and cultivated feedstock needs more inputs (especially fertilizers and pesticides) to be produced [3] so this result was unexpected. However, it is also well known that perennial energy crops can stock carbon underground [91]. Therefore, our counter-intuitive result can be explained by this fact, but only if direct LUC is accounted for (accounting for above ground and underground carbon sequestration). However we noticed that *luc\_dir* variable is not statistically significant. Hence, we decided to combine *mat\_cult* variable with *luc\_dir* variable (aggregated in *mat\_cultxdluc*) in order to confirm this effect (results reported in columns (1aG2) and (2aG2), Table 4). Our meta-model shows that *mat\_cultxdluc* variable is statistically significant at the 1% level and impacts negatively GHG emissions for G2 biofuels. It means that GHG emissions for G2 biofuels produced from cultivated feedstock that take into account dLUC are lower than GHG emissions for G2 biofuels from cultivated feedstock that do not take into account dLUC or from waste feedstock. Thus, the type of feedstock combined with the fact to take into account dLUC influence GHG emissions for G2 biofuels.

#### Methodological variables

*lca\_cons* variable is statistically significant at the 1% level for "G2" sample. Its coefficient is negative so GHG emissions for G2 biofuels are lower with a consequential approach compared to the attributional approach. The type of LCA approach thus influences GHG emission results for G2 biofuels.

*copval\_alloc* and *copval\_hyb* variables are statistically significant at the 1% and 5% level, respectively (column (1aG2), Table 4). It confirms the influence of the method for taking into account coproducts on LCA GHG emission results as often mentioned in the literature [60]. The coefficients of both variables are positive which means that GHG emissions are lower for G2 biofuels when using the system boundaries expansion method (*copval\_systexp*) compared to allocation method and hybrid method. We observed, however, that most LCA authors recognize the importance of the method of taking into account burdens associated to coproducts. 91% of the studies in our database test alternative methods for allocation performing a sensibility analysis.

*luc\_indir* is statistically significant at the 1% level. It shall be noticed that all studies assessing indirect LUC (*luc\_indir*) always assess direct LUC (*luc\_dir*), so *luc\_indir* is equal to 1 when the study assesses both direct and indirect LUC. Nevertheless *luc\_dir* is not statistically significant. We can then conclude that assessing indirect LUC increases GHG emission results for G2 biofuels as *luc\_indir* coefficient is positive. Nevertheless, the direct LUC (*luc\_dir*) has an influence but it is linked with the type of biomass feedstock used, as mentioned before.

*impcat\_nev*, *impcat\_nrc* variables are both statistically significant at the 1% level. The type of other environmental indicators than GHG emissions assessed in the study thus could influence GHG emission results for G2 biofuels. According to our results, GHG emissions are statistically lower when the study assesses the Net Energy Value (*impcat\_nev*) and are statistically higher when the study assesses the Non Renewable Energy consumption (*impcat\_nrc*). This effect could not have been anticipated. Nevertheless, it could be interpreted as a quality indicator for the study: when these energy indicators are consistently assessed, the GHG emission result can be considered to be more robust.

Variables related to the methods for taking into account uncertainties (*uncer\_MC* and *uncer\_SA*) are statistically significant and impact positively the amount of GHG emissions emitted for G2 biofuels. This effect is unexpected. It means that GHG emissions for G2 biofuels are statistically higher when uncertainties are taken into account – via Monte Carlo method (*uncer\_MC*) or Sensitivity analysis (*uncer\_SA*) – than when there is no uncertainties assessment (*uncer\_ref*). The fact to assess uncertainties could be interpreted also as a quality indicator of a study. It can be seen as an effort to establish the accuracy of the results but the tendency of the influence of these parameters in the e-s could not be anticipated nor explained afterwards.

### Typological variables

Lastly, *zlab\_us* variable has a negative impact on GHG emissions and is significant at the 1% level. Again, GHG emissions for G2 appear to be statistically lower when the authors are from North America (*zlab\_us*) compared to authors from Europe (*zlab\_eu*). Hence, the geographical location of the authors also influences GHG emission results for G2 biofuels.

### **4.2.3 Results for the Ethanol sample**

Estimates results for the "G2-Ethanol" sample are presented in Table 5. Columns (1cEtha) and (2cEtha) correspond to the model without the inclusion of the technical variable that is the mass yield of the pathway (*g2\_mass\_yield*). Columns (1bEtha) and (2bEtha) test the existence of a linear effect of this variable (*g2\_mass\_yield*) whereas columns (1aEtha) and (2aEtha) test the existence of a non-linear effect of this variable by taking the logarithm of the *g2\_mass\_yield* variable (*g2\_mass\_yield\_ln*). The AIC and the BIC both decrease from the first specification (columns (1aEtha) and (2aEtha)) to the last one (columns (1cEtha) and (2cEtha)). Therefore, the inclusion of a non-linear effect of the mass yield of the pathway appears more relevant to explain GHG emission variations. Thus, we choose to comment results presented in column (1aEtha).

### Technical variables

*mat\_cultxdluc* variable is significant at the 5% level and has the same effect on GHG emissions for Ethanol as for G2 biofuels (See Section 4.2.2)

The mass yield of the pathway *g2\_mass\_yield\_ln* impacts negatively GHG emissions for G2 Ethanol, which is an intuitive effect: the better the mass yield is, the less GHG are emitted all along the biofuel life cycle, *ceteris paribus*. It should be noticed that *g2\_mass\_yield\_ln* traduces a non-linear effect of this variable.

We should also mention that variables related to other technical data, such as the type of biomass pretreatment, are not statistically significant for Ethanol. Indeed, 83% of observations are related to Ethanol produced using dilute sulfuric acid pretreatment and most of these observations use technical data from the same study (NREL) [92]. Hence pretreatment process variables for Ethanol are not really discriminatory, and this could explain why those variables are not statistically significant.

### Methodological variables

Among significant variables found for G2 biofuel sample, *lca\_cons*, *luc\_indir*, *impcat\_nev*, *impcat\_nrc*, *uncer\_MC* and *uncer\_SA* variables are also significant for the Ethanol sample and have the same impact as described for the G2 sample. So the type of LCA approach, the fact 'to assess indirect LUC', the type of other environmental indicators, the method for taking into account uncertainties influence GHG emission results for G2 Ethanol.

It can be noticed that *copval\_alloc* and *copval\_sys* variables are no longer statistically significant. This result is surprising regarding a previous lignocellulosic Ethanol LCA studies review [13] which concludes that the treatment of coproducts has a strong influence in the LCA results.

### Typological variables

*zlab\_us* variable has a negative impact on GHG emissions and is significant at the 5% level. It means that GHG emissions for Ethanol are statistically lower when the authors are from North America (*zlab\_us*) compared to authors from Europe (*zlab\_eu*). Hence, the geographical location of the authors also influences GHG emission results for Ethanol.

**Table 5 – Results of MRA for the econometric samples G2-Ethanol biofuels**

Samples Model	Ethanol 1aEtha	Ethanol 2aEtha	Ethanol 1bEtha	Ethanol 2bEtha	Ethanol 1cEtha	Ethanol 2cEtha
Constant	-5.88 (14.8)	31.9 (31.58)	46.39*** (8.33)	37.85*** (13.52)	32.11*** (2.64)	34.07*** (3.95)
<b>Technical data</b>						
mat_cultxdluc	-7.18** (3.26)	-10.1** (4.85)	-6.91** (3.28)	-9.92** (4.79)	-	-
g2_mass_yield			-73.57** (34.79)	-		
g2_mass_yield_sq						
g2_mass_yield_ln	-23.22** (9.3)	-				
<b>Methodological choices</b>						
lca_att (ref)						
lca_cons	-38.24*** (5.53)	-40.12*** (11.37)	-38.43*** (5.46)	-40.17*** (11.33)	-40.24*** (4.72)	-40.72*** (9.9)
luc_indir	29.01*** (7.49)	35.89*** (8.59)	27.48*** (7.41)	35.07*** (8.43)	19.85*** (7.29)	31.7*** (7.78)
uncer_MC	9.51** (4.06)	20.32*** (5.83)	9.66** (4.11)	20.43*** (5.82)	10.06** (4.23)	18.67*** (5.5)
uncer_SA	12.5*** (3.68)	12.28** (5.71)	11.53*** (3.59)	11.99** (5.57)	12.09*** (2.72)	14.01*** (4.08)
uncer_ref (ref)						
impcat_nev	12.34*** (3.99)	-	11.05*** (3.99)	-	9.51** (3.94)	11.15* (6.49)
impcat_nrc	-17.09*** (3.19)	-14.16*** (5)	-17.24*** (3.17)	-14.18*** (5)	-22.53*** (2.51)	-20.14*** (4.08)
impcat_other	-	-	-	-	-1.09* (0.61)	-
impcat_gwponly (ref)						
<b>Typology of the study</b>						
zlab_us	-7.29** (3.56)	-28.56*** (7.44)	-8.22** (3.56)	-29.23*** (7.35)	-11.26*** (2.37)	-23.84*** (4.74)
zlab_eu (ref)						
<b>Model information</b>						
N	209	209	209	209	321	321
Mean dep. Var.	19.70	19.14	19.70	18.96	26.61	26.61
Adj. R-squ.	31.21%	36.31%	30.32%	36.24%	40.13%	37.82%
Log-Likelihood	-884.15	-919.42	-885.49	-919.54	-1364.10	-1412.35
F-stat.	12.61	10.91	12.55	10.79	33.94	27.19
(P. value)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)
Skewness	18.14		16.84		18.16	
(P. value)	(0.0527)		(0.078)		(0.0333)	
Kurtosis	0.32		0.23			
(P. value)	(0.5729)		(0.6351)			
AIC	1790.30	1860.83	1792.98	1861.08	2748.21	2844.70
BIC	1827.07	1897.60	1829.75	1897.85	2785.92	2882.42
LR test (P. value)	959.9	985.87	957.22	985.62		
Nested model: model (c)	(0,0000)	(0,0000)	(0,0000)	(0,0000)		
Procedure	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS

#### 4.2.4 Results for the BtL sample

Estimates results for the "G2-BtL" sample are presented in Table 6. Columns (1eBtL) and (2eBtL) correspond to the reduced model obtained for the "G2" sample. Columns (1dBtL) and (2dBtL) correspond to the *new* reduced model without any technical variables. Columns (1aBtL) to (2cBtL) correspond to the reduced model with technical variables. Columns (1aBtL) and (2aBtL) are the only one to test a non-linear effect of the mass yield of the pathway. The AIC and the BIC both decrease from the first specification (columns (1aBtL) and (2aBtL)) to the last one (columns (1eBtL) and (2eBtL)). Thus, we choose to comment results presented in column (1aBtL).

##### Technical variables

*mat\_cultxdluc* variable is significant at the 1% level and has the same effect on GHG emissions for BtL as for G2 biofuels (See Section 4.2.2)

Variables related to the type of conversion into fuel process (*btl\_pro\_alng* and *btl\_pro\_alelec*) are statistically significant. Using natural gas as a source of heat for an allothermic BtL unit leads to higher GHG emissions than producing BtL from an autothermic plant (biomass provides all process energy needs). Conversely using grid electricity as a utility for an allothermic BtL unit leads to lower GHG emissions than producing BtL from an autothermic plant. The source of electricity used could explain these results. Indeed, among the observations using grid electricity as a utility for an allothermic BtL unit, 57% of these observations use electricity provided by wind power plants [58]. The other studies do not precise the source of electricity used.

The mass yield of the pathway *g2\_mass\_yield\_ln* impacts negatively GHG emissions for BtL, which is an expected effect: the better the mass yield is, the less GHG emissions are emitted all along the pathway for a G2 biofuel. It should also be noticed that *g2\_mass\_yield\_ln* traduces a non-linear effect of this variable.

It should be noticed that variables related to other technical data, such as the type of biomass pretreatment or the inclusion of Carbon Capture and Storage (CCS) in the process, are not statistically significant for BtL. Indeed, 90% of observations in the econometric sample are related to BtL produced without biomass pretreatment (See Table C.4 in Appendix C). Hence pretreatment process variables for BtL are not really discriminatory, and this may be the reason why those variables are not statistically significant. Moreover, the variable *btl\_ccs* is equal to zero for the econometric sample (See Table C.4 in Appendix C), therefore this variable could not have been tested. In fact, the variable in question appears in only three observations and all of them are considered outliers (see Table B.8 in Appendix B).

#### Methodological variables

Among significant variables found for "G2" sample, only *copval\_alloc* and *lca\_cons* are significant for the "G2-BtL" sample. The method for taking into account coproducts (*copval\_alloc*) has the same impact as described for "G2" sample (*copval\_hyb* for BtL is equal to zero). However the influence of the type of LCA approach is not the same for G2 biofuel and for BtL: GHG emissions are higher with a consequential approach (*lca\_cons*) compared to an attributional approach (*lca\_att*). So the type of LCA approach and the method for taking into account coproducts influence GHG emission results for BtL.

Furthermore, the type of coproduct influence GHG emission results for BtL since *cop\_elec* variable is statistically significant at the 1% level. So the coproduction of electricity for BtL decreases GHG emissions compared to other coproducts, *ceteris paribus*.

#### Typological variables

*zlab\_us* variable has a negative impact on GHG emissions and is significant at the 1% level. It means that GHG emissions for BtL are statistically lower when the authors are from North America (*zlab\_us*) compared to authors from Europe (*zlab\_eu*). Hence, the geographical location of the authors also influences GHG emission results for BtL.

**Table 6 – Results of MRA for the econometric samples G2-BtL biofuels**

Samples Model	BtL	BtL								
	1aBtL	2aBtL	1bBtL	2bBtL	1cBtL	2cBtL	1dBtL	2dBtL	1eBtL	2eBtL
Constant	43.23*** (12.66)	29.68 (20.05)	70.94*** (9.38)	75.11*** (12.85)	39.6*** (6.29)	32.19*** (6.36)	27.43*** (5.59)	29.03*** (5.69)	16.03** (6.33)	24.5*** (8.38)
<b>Technical data</b>										
mat_cultxdluc	-14.41*** (3.08)	-19.53*** (3.27)	-15.41*** (3.11)	-20.33*** (3.23)	-15.16*** (2.93)	-18.1*** (2.84)	-16.24*** (2.47)	-17.02*** (2.6)	-11.64*** (3.98)	-13.28*** (4.46)
cop_elec	-44.56*** (4.14)	-35.68*** (7.7)	-43.99*** (4.19)	-35.94*** (7.71)	-19.4*** (5.09)	-	-	-	-	-
g2_mass_yield	-	-	-	-	-	-	-	-	-	-
g2_mass_yield_sq	-	-	-	-	-	-	-	-	-	-
g2_mass_yield_ln	-11.66* (7.04)	-17.41** (8.52)	-	-	-	-	-	-	-	-
btl_pro_autoth	-	-	-	-	-	-	-	-	-	-
btl_pro_alng	17.04*** (5.4)	-	17.73*** (5.73)	-	13.85** (5.6)	-	-	-	-	-
btl_pro_alelec	-	-	-	-	-	-	-	-	-	-
btl_pro_alrenew	-	-	-	-	-	-	-	-	-	-
btl_gasrecycl	-	-	-	-	-	-	-	-	-	-
<b>Methodological choices</b>										
lca_att (ref)	-	-	-	-	-	-	-	-	-	-
lca_cons	25.61*** (8.36)	-	26.23*** (8.6)	-	22.97*** (8.68)	-	18.1** (8.87)	-	-	-
copval_alloc	9.6*** (3.37)	-	9.81*** (3.4)	-	13.73*** (3.05)	12.51*** (4.33)	16*** (2.96)	11.02** (4.65)	11.7*** (2.79)	13.21*** (4.31)
copval_systexp (ref)	-	-	-	-	-	-	-	-	-	-
copval_hyb	-	-	-	-	-	-	-	-	-	-
luc_indir	-	-	-	-	-	-	-	-	-	-
uncer_MC	-	-	-	-	-	-	-	-	-	-
uncer_SA	-	-	-	-	-	-	-	-	-	-
uncer_ref (ref)	-	-	-	-	-	-	-	-	-	-
impcat_nev	-	-	-	-	-	-	-	-	-	-
impcat_nrc	-	-	-	-	-	-	-	-	-	-
impcat_other	-	-	-	-	-	-	-	-	-	-
impcat_gwponly (ref)	-	-	-	-	-	-	-	-	-	-
<b>Typology of the study</b>										
zlab_us	-21.42*** (4.8)	-	-24.38*** (4.71)	-16.37* (9.57)	-22.11*** (4.35)	-16.59*** (5.51)	-16.51*** (4.09)	-14.91*** (5.03)	-11.41** (4.93)	-17.34*** (5.29)
zlab_eu (ref)	-	-	-	-	-	-	-	-	-	-
<b>Model information</b>										
N	132	132	132	132	141	141	143	143	143	143
Mean dep. Var.	19.45	21.96	19.45	21.84	18.80	22.29	18.65	22.22	18.65	21.62
Adj. R-squ.	39.48%	26.01%	38.39%	23.56%	35.19%	25.06%	31.39%	25.56%	33.22%	24.68%
Log-Likelihood	-548.53	-568.94	-549.72	-571.09	-589.14	-608.32	-603.08	-618.57	-599.04	-617.29
F-stat.	-	-	-	-	-	-	16	15.66	11.34	10.69
(P. value)	-	-	-	-	-	-	(0,0000)	(0,0000)	(0,0000)	(0,0000)
Skewness	12.64	-	13.7	-	13.31	-	12.09	-	16.91	-
(P. value)	(0.2445)	-	(0.1869)	-	(0.1492)	-	(0.0336)	-	(0.0501)	-
Kurtosis	2.53	-	2.38	-	2.22	-	1.64	-	1.52	-
(P. value)	(0.1117)	-	(0.1232)	-	(0.1364)	-	(0.2004)	-	(0.2184)	-
AIC	1115.07	1155.89	1117.44	1160.18	1194.29	1232.65	1218.17	1249.13	1218.07	1254.57
BIC	1141.01	1181.83	1143.38	1186.12	1217.88	1256.24	1235.95	1266.91	1247.70	1284.20
LR test (P. value)	109.1	99.25	106.73	94.96	27.88	20.49	-	-	8.09	2.56
Nested model: model (d)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	-	-	(0,0882)	(0,6336)
Procedure	OLS (White's HCCM)	WLS								

We turn now to the analysis of the "G3" sample.

#### 4.2.5 Results for the G3 sample

Estimate results for the "G3" sample are presented in Table 7. We begin by commenting the impact of *g3\_productivity* and *g3\_oil* as the influence of these two continuous technical variables will determine the final specification of the model for the "G3" sample.

##### Technical variables

First, a lin-lin model is specified in order to test the linear effects of both *g3\_productivity* and *g3\_oil* on the *e-s*. Table 7, columns (1dG3) and shows the reduced form of this specification. It could be noticed that *g3\_productivity* variable is not statistically significant. This result is non intuitive as most part of the literature mentions that algae productivity could explain the variability of GHG emission results. The non-significance of this variable may be explained by the existence of a non-linear effect instead of a linear one. To test this hypothesis, two models are specified. In the first one (Table 7, column (1cG3)), the non-linear effect is modelled as a second-degree polynomial by introducing the variable *g3\_productivity* and its

squared value ( $g3\_productivity\_sq$ ). In the second one (Table 7, column (1bG3)), the linear effect is modelled as a logarithmic function by introducing  $g3\_productivity\_ln$  instead of  $g3\_productivity$ . In Table 7, column (1cG3), neither  $g3\_productivity$  nor  $g3\_productivity\_sq$  are statistically significant at the 10% level. On the contrary,  $g3\_productivity\_ln$  is statistically significant at the 1% level (Table 7, column (1bG3)). As a conclusion, the variable  $g3\_productivity$  does have an impact on GHG emission results for G3 biofuels but this effect is non-linear which can be captured by a logarithmic function, not a second degree polynomial. Regarding  $g3\_oil$ , results presented in Table 7, column (1bG3) indicates a negative linear influence of this variable. Finally only  $g3\_productivity\_ln$  and  $g3\_oil\_ln$  variables are statistically significant at the 1% level and their coefficients are both negative (Table 7, column (1aG3)). Thus, we choose to comment results presented in column (1aG3).

Algae productivity value and the oil content – as proxies of the  $g3\_productivity$  and  $g3\_oil$  variables, respectively – influence GHG emission results for G3 biofuels. They have a negative impact on GHG emissions so the higher algae productivity or the algae content is, the lower GHG emissions are. In addition, these non-linear effects indicate that those parameters are more sensitive for low productivity or low oil content than for high ones.

The variable  $hvo$  is statistically significant at the 1% level. According to its coefficient estimate, GHG emissions for HVO from algae are higher than GHG emissions from FAME from algae by about 134 g CO<sub>2</sub>eq/MJ *ceteris paribus*. It indicates that the type of fuel conversion technology can explain the variability of GHG emission results for G3 biofuels. This result is difficult to be interpreted, especially due to the extent of its coefficient. In fact, the literature shows that upstream fossil energy consumption (including all inputs, notably methanol and hydrogen production) is similar in FAME and HVO processes [93]<sup>15</sup>. Vegetable oil consumption for both processes is also quite similar.

The coefficient of  $g3\_Oppond$  is negative and significant at the 1% level. GHG emissions for G3 biofuels are thus statistically lower when microalgae are grown in an open-pond than in a photobioreactor. Hence the type of technology used for microalgae cultivation influences GHG emission results for G3 biofuels.

The type of technology used for microalgae cultivation, the algae productivity and the oil content of algae are often identified as key parameters in G3 biofuel LCA studies. So the fact that those variables are statistically significant confirms previous conclusions found in the literature. Jorquera *et al.* [94], in a microalgae LCA study (not included in this review because conversion into biofuel is not included), shows that culture in photobioreactors is more energy intensive than in open ponds. One of the conclusions of previous literature review on biofuel from microalgae technologies [95,96] is that microalgae strains presenting high biomass productivity are better for CO<sub>2</sub> emission mitigation.

#### Methodological variables

Concerning methodological variables, only  $lca\_cons$  variable is statistically significant at the 1% level for G3 sample. Its positive coefficient indicates that GHG emissions for G3 biofuels are statistically higher when the study uses a consequential approach for LCA compared to the attributional approach. Hence the type of LCA approach influences GHG emission results for G3 biofuels. However, note that consequential LCA approach is only used by one study (that represents 9% of the observations for the econometric sample). Consequently the influence of the type of LCA approach for G3 biofuels should be interpreted with caution.

Liu *et al.* [17], in an LCA harmonization exercise, show that different authors accounted for different microalgae coproducts and that it plays an important role in the final life cycle GHG emissions of the biofuel. However, in our meta-regression, variables related to the coproducts did not show themselves to be

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<sup>15</sup> In the EcoInvent database [116], the cumulative fossil energy demand for the production of 1 kg of hydrogen from cracking natural gas is 70, 9 MJ. The same indicator for 1 kg of methanol also produced from natural gas is 36.9 MJ. FAME contains around 10% of methanol and HVO around 4% of hydrogen (mass). The selected processes for this example are the most commonly used for these products.

statistically significant. Still, the fact that *lca\_cons* is statistically significant warns us about the importance of the definition of system boundaries and coproduct accounting methodology.

### Typological variables

Regarding typological variables, the coefficient of *zlab\_us* variable is significant at the 1% level and its sign is negative whereas *zlab\_other* is not statistically significant. Thus, the previous result regarding the influence of geographical location is partly retrieved: GHG emissions of G3 biofuels are statistically lower when studies are from NA compared to ones from Europe. The non-significance of *zlab\_other* indicates that there are no systematic differences between results drawn from European studies and other countries.

**Table 7 – Results of MRA for the econometric samples G3 biofuels**

Samples Model	G3 1aG3	G3 2aG3	G3 1bG3	G3 2bG3	G3 1cG3	G3 2cG3	G3 1dG3	G3 2dG3	G3 1eG3	G3 2eG3
Constant	318.44*** (90.09)	550.41*** (171.08)	621.72*** (99.61)	916.55*** (146.59)	490.82*** (87.74)	640.23*** (68.43)	450.73*** (88.11)	585.43*** (59.96)	105.38*** (19.91)	237.46*** (25.17)
<b>Technical data</b>										
fame										
hvo	134.18*** (35.34)	185.12*** (39.47)	135.18*** (32.44)	181.47*** (34.94)	137.73*** (34.09)	178.64*** (35.28)	134.31*** (34.62)	176.78*** (34.69)		
g3_productivity					-	-5.82*** (1.86)	-	-3.19*** (1.2)		
g3_productivity_sq					-	0.02** (0.01)				
g3_productivity_ln	-65.31*** (20.13)	-124.8*** (45.43)	-64.46*** (20.06)	-127.33*** (44.5)						
g3_oil			-434.74*** (142.4)	-521.06*** (112.68)	-425.28*** (150.9)	-522.41*** (110.74)	-430.6*** (146.08)	-527*** (115.09)		
g3_oil_sq										
g3_oil_ln	-140.32*** (42.3)	-161.66*** (39.17)								
g3_Oppond	-197.13*** (34.6)	-259.9*** (24.28)	-198.94*** (36.92)	-260.33*** (24.35)	-201.93*** (38.32)	-257.79*** (24.13)	-199.6*** (38.46)	-257.1*** (23.73)		
<b>Methodological choices</b>										
lca_att (ref)										
lca_cons	172.72*** (61.08)	250.7*** (70.3)	174.8*** (65.98)	254.66*** (73.36)	196.92** (79.77)	268.68*** (81.63)	187.41** (84.67)	290.31*** (86.21)	-	-
<b>Typology of the study</b>										
zlab_us	-207.56*** (31.01)	-259.02*** (26.35)	-198.73*** (29.55)	-244.44*** (25.73)	-201.53*** (30.19)	-244.11*** (26.09)	-199.27*** (30.8)	-240.76*** (25.61)	-95.93*** (26.17)	-225.96*** (31.21)
zlab_eu (ref)										
zlab_other	-	-	-	-	-	-	-	-	-114.8*** (21.87)	-246.87*** (26.35)
<b>Model information</b>										
N	68	68	68	68	68	68	68	68	69	69
Mean dep. Var.	59.97	68.95	59.97	67.95	59.97	67.85	59.97	66.53	58.84	126.49
Adj. R-squ.	65.23%	80.63%	66.07%	81.32%	66.59%	81.92%	66.06%	81.34%	17.77%	48.39%
Log-Likelihood	-373.01	-376.38	-372.18	-375.14	-371.08	-373.46	-372.19	-375.11	-410.22	-418.31
F-stat.	11.11	24.7	11.67	25.07	10.96	22.62	13.53	27.17	11.17	31.42
(P. value)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)	(0,0000)
Skewness	9.25		13.93		14.15		16.85		32.57	
(P. value)	(0.2352)		(0.0524)		(0.078)		(0.0184)		(0,0000)	
Kurtosis	0.06		0.32		0.47		0.41		6.01	
(P. value)	(0.8139)		(0.5694)		(0.4938)		(0.521)		(0.0142)	
AIC	762.02	768.75	760.36	766.29	760.17	764.92	760.37	766.23	828.44	844.62
BIC	779.78	786.51	778.11	784.04	780.14	784.89	778.13	783.98	837.37	853.56
LR test (P. value)	74.42	83.87	76.08	86.34	78.27	89.71	76.06	86.4		
Nested model: model (e)	(0,0000)	(0,0000)	(0,0000)	(0)	(0,0000)	(0,0000)	(0,0000)	(0,0000)		
Procedure	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS	OLS (White's HCCM)	WLS

## 4.3 Harmonization

The MRA results presented in Section 4.2 are now used to address the harmonization issue in the field of advanced biofuels GHG emissions thanks to the technique of *benefits transfer using meta-regression models*. As already demonstrated in the previous Section, the *meta-regression* framework allows the production of an estimation of the mean *e-s* weighted by the systematic influence of its main drivers. Once estimated, the meta-function can be used to deduce original values of the *e-s* by specifying new values for the main drivers identified corresponding to relevant case studies. This technique of *benefits transfer using*

*meta-regression models*, as it is named in the *MA* literature, may be a particularly well adapted methodology to deal with the so-called harmonization issue specific to the LCA literature.

This section aims at providing an illustration of the potential for *MRA* to perform harmonization in the field of LCA through an application to advanced biofuels GHG emissions. To do so, predicted values of the *e-s* are computed using the meta-functions estimated in Section 4.2.

The predicted values can be calculated using a combination of variables that already exists in the meta-database: this type of prediction is called "in sample". In sample prediction enables the comparison of collected values (estimations of the *e-s*) and predicted values in order to check the accuracy of the meta-function in predicting the *e-s*.

Furthermore, predicted values can be extrapolated for a combination of relevant variables that do not necessarily exist in the meta-database, hence the prediction is called "out of sample". Out of sample prediction could provide values for the *e-s* for case studies not assessed in the literature. In addition, out of sample prediction applied to quantitative variables can help to test how sensible the *e-s* is to these variables.

First, in sample predictions are presented and analyzed. Second, out of sample predictions are conducted assessing in particular the sensitivity of quantitative variables (algae productivity and oil content for G3 biofuels, mass yield for BtL and Ethanol).

### 4.3.1 Prediction in sample

Table 8 presents some characteristics of predicted values compared to collected values (estimations of the *e-s* in the meta-database) for each sample. The meta-models used to calculate these *in sample predictions* are those estimated in columns (1aAll), (1aG2), (1aEth), (1aBtL) and (1aG3) for the "whole" sample, the "G2" sample, the "G2-Ethanol" sample, the "G2-BtL" sample and the "G3" sample, respectively (see Table 4, Table 5, Table 6 and Table 7).

First, we observe that mean values for predicted values are slightly different from those of collected values. Nevertheless the ranking between G2 and G3 biofuels, BtL and Ethanol in terms of contribution to the climate change (i.e. amount of GHG emissions emitted all along their life cycle) is still the same as depicted in the econometric analysis. Second, the range of variation is narrower for predicted values than for collected values, except for the G3 sample. Furthermore, these meta-models tend to overestimate predicted values compared to their corresponding collected values (53% to 56% of predicted values are overestimated depending on the samples) as depicted in Figure 6.

**Table 8 – Characteristics of collected and predicted values of the *e-s* in g CO<sub>2</sub>eq/MJ (predicted values calculated from (1a) meta-models)**

Samples	Whole	G3	G2	BtL	Ethanol
<b>Collected values</b>					
Number of values	533	69	464	143	321
Mean	28.64	58.84	24.15	18.65	26.61
Min	-85.00	-85.00	-24.00	-24.00	-23.65
Max	332.20	332.20	85.80	85.68	85.80
NA values higher than -60% GHG emission threshold	7%	14%	5%	1%	7%
EU values higher than -60% GHG emission threshold	25%	30%	24%	17%	27%
<b>Predicted values</b>					
Number of values	533	68	464	132	209
Mean	28.64	59.97	24.15	19.45	19.7
[Confident Interval]	[25.19;32.09]	[43.29;76.65]	[22.56;25.74]	[16.67;22.23]	[17.37;22.03]
Min	-9.42	-109.25	-15.82	-8.04	-20.86
Max	76.27	230.82	47.91	56.31	47.49
Underestimated values	44%	46%	44%	47%	47%
Overestimated values	56%	54%	56%	53%	53%
Collected values included in the predicted value CI	12%	51%	22%	18%	37%
NA values higher than -60% GHG emission threshold	5%	9%	2%	1%	2%
EU values higher than -60% GHG emission threshold	7%	28%	22%	8%	1%

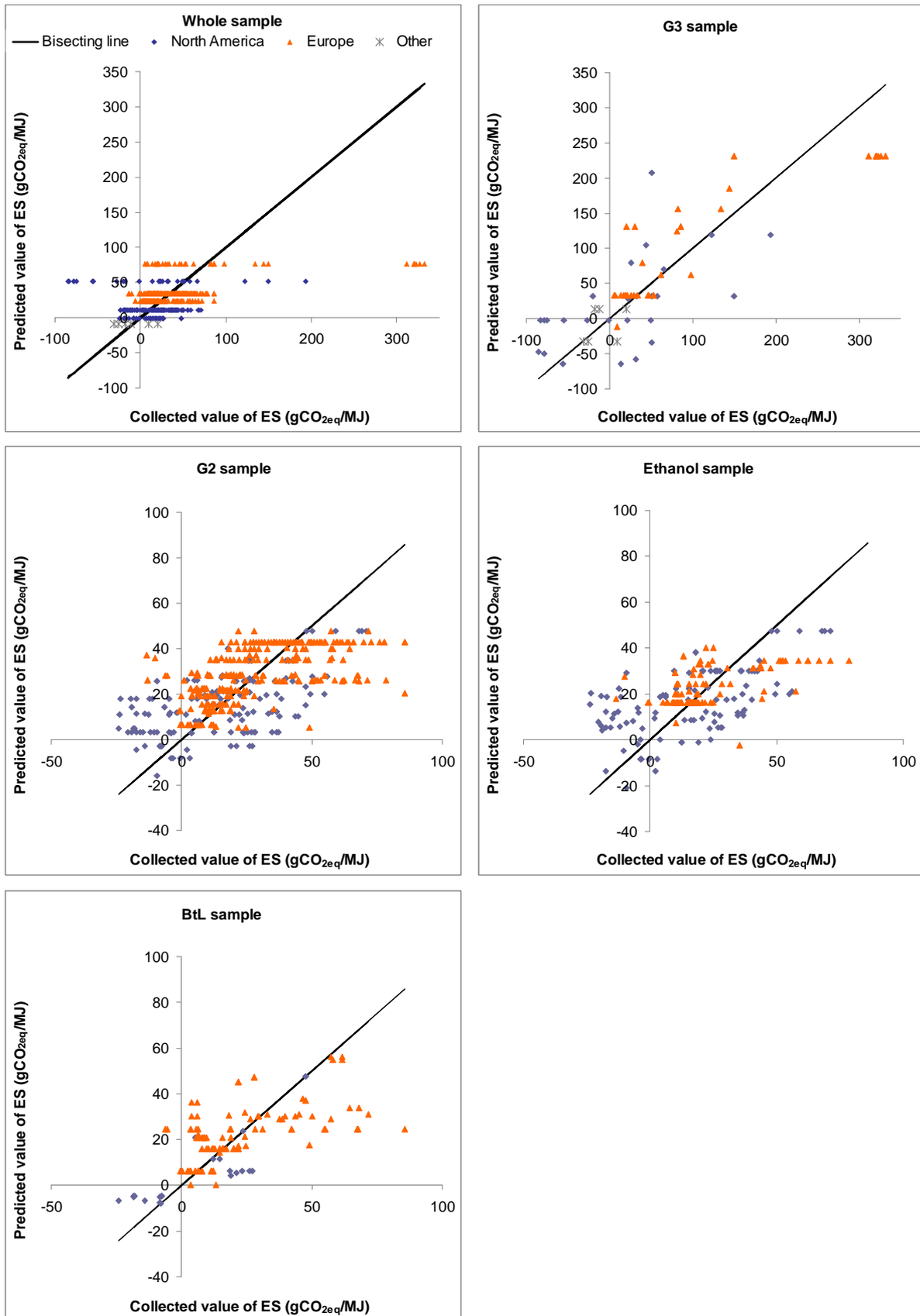


Figure 6 – Predicted and collected values of the  $e$ -s for meta-model (1a) distinguished by their geographical location

### 4.3.2 Prediction out of sample

Out of sample prediction enables the building of values of the  $e$ - $s$  for combinations of variables that do not necessarily exist in the meta-database. Those values are calculated from the meta-function obtained by the meta-regression method. This harmonization method allows us to obtain mean values of the  $e$ - $s$  and associated confidence intervals (CI) for each combination of statistically significant variables of a meta-model. For instance, using the meta-model for the "whole" sample presented in column (1aAll), Table 4, predicted values of the  $e$ - $s$  can be calculated for G3 biofuel, BtL and Ethanol in Europe and North America. Table 9 and Table 10 illustrate the procedure. Table 9 reports coefficient estimates of the model (1aAll) (as presented in column (1aAll), Table 4) and the different values of the variable of this reduced model which have to be imputed to compute the predicted values of the  $e$ - $s$  for G3 biofuel, BtL and Ethanol in Europe and North America. Table 10 shows the link between these imputed values and the corresponding predicted values of the  $e$ - $s$  whereas Figure 7 offers an alternative view of Table 10 results.

**Table 9 - Benefits Transfer for the "Whole" Sample (1aAll meta-model)**

Samples Model: Parameter Estimate	Whole 1aAll	Imputed Values					
Constant	76,27*** (13,64)	1	1	1	1	1	1
<b>Technical data</b>							
gen_3 (ref for Whole)		0	0	0	0	0	0
etha	-41,39*** (13,14)	0	1	0	0	1	0
btl (ref for G2)	-52,12*** (13,36)	0	0	1	0	0	1
<b>Typology of the study</b>							
zlab_us	-24,6*** (3,97)	0	0	0	1	1	1
zlab_eu (ref)		0	0	0	0	0	0
zlab_other	-85,69*** (15,6)	0	0	0	0	0	0
<b>Transfer Values</b>		<b>76,27</b> <b>(13,64)</b>	<b>34,88</b> <b>(1,75)</b>	<b>24,15</b> <b>(1,88)</b>	<b>51,67</b> <b>(12,35)</b>	<b>10,29</b> <b>(2,98)</b>	<b>-0,44</b> <b>(3,50)</b>

**Table 10 - Harmonized  $e$ - $s$  (g CO<sub>2</sub>eq/MJ) for the "Whole" sample (1aAll meta-model))**

		Harmonized $e$ - $s$	95% Confidence Interval	
			Min	Max
Europe	G3	76,27	49,54	103,00
	G2 Ethanol	34,88	31,45	38,31
	G2 BtL	24,15	20,46	27,84
North America	G3	51,67	27,47	75,88
	G2 Ethanol	10,29	4,45	16,13
	G2 BtL	-0,44	-7,31	6,42
<b>In sample predicted value</b>	<b>Mean</b>	<b>28,64</b>	<b>25,19</b>	<b>32,09</b>

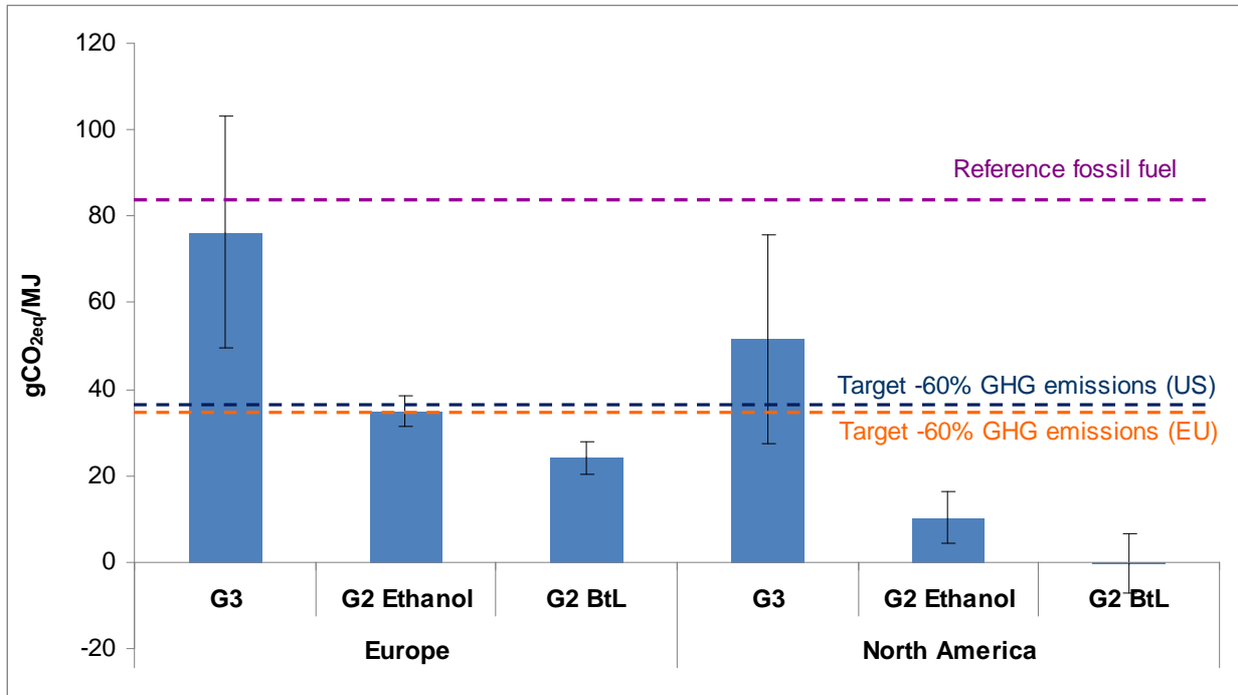


Figure 7 – Predicted values of the effect size for the whole sample calculated from meta-model 1aAll

As depicted in the Figure 7, predicted values of GHG emissions for advanced biofuels in Europe are always higher than those in North America. In addition, GHG emissions are lower for BtL than Ethanol, and G3 biofuels always emit more GHG emissions than G2 biofuels. Those results are in line with the statistical description conducted in Section 3.2. Furthermore, the predicted value CIs are wider for G3 biofuels than for G2 biofuels, meaning that the model better estimates G2 biofuels GHG emissions than those of G3 biofuels. It should be noted that predicted values of GHG emissions for advanced biofuels are always lower than GHG emissions for the reference fossil fuel even when considering CI, except for G3 biofuel in Europe.

The same type of analysis could be conducted for each meta-model. Out of sample prediction could also be used to test the sensitivity of results for quantitative variables. A range of values for quantitative variables could be tested by calculating mean predicted values for the *e-s* and associated CI, *ceteris paribus*.

For instance, the influence of oil content and algae productivity is tested for G3 biofuels (Figure 8 and Figure 9), by testing the range of values found in the meta-database. Results show that both variables have a non-linear effect on LCA GHG emissions, *ceteris paribus*. Furthermore, variations for high values of the algae productivity have less effect on the *e-s* than variations for low values. Moreover, CIs are smaller for oil content and algae productivity values around mean values than for extreme values.

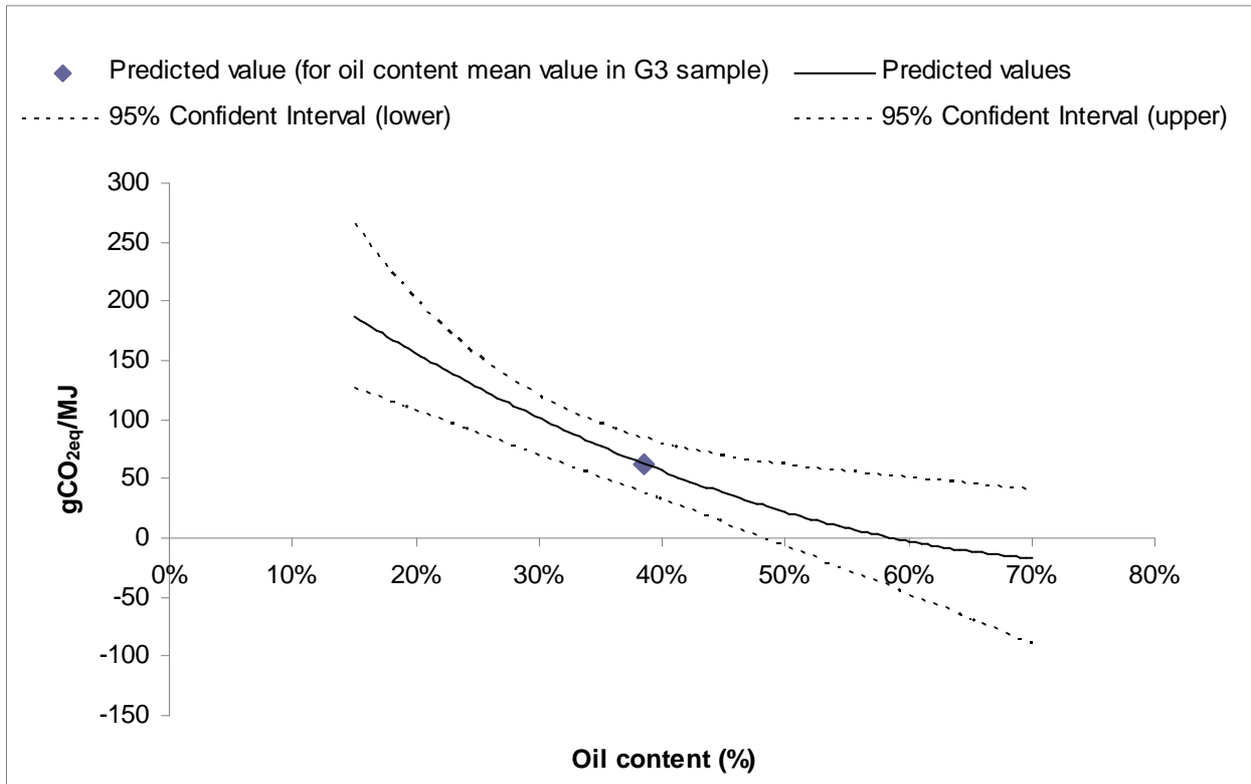


Figure 8 – Influence of oil content on predicted values of the  $e$ -s for G3 sample (1aG3 meta-model)

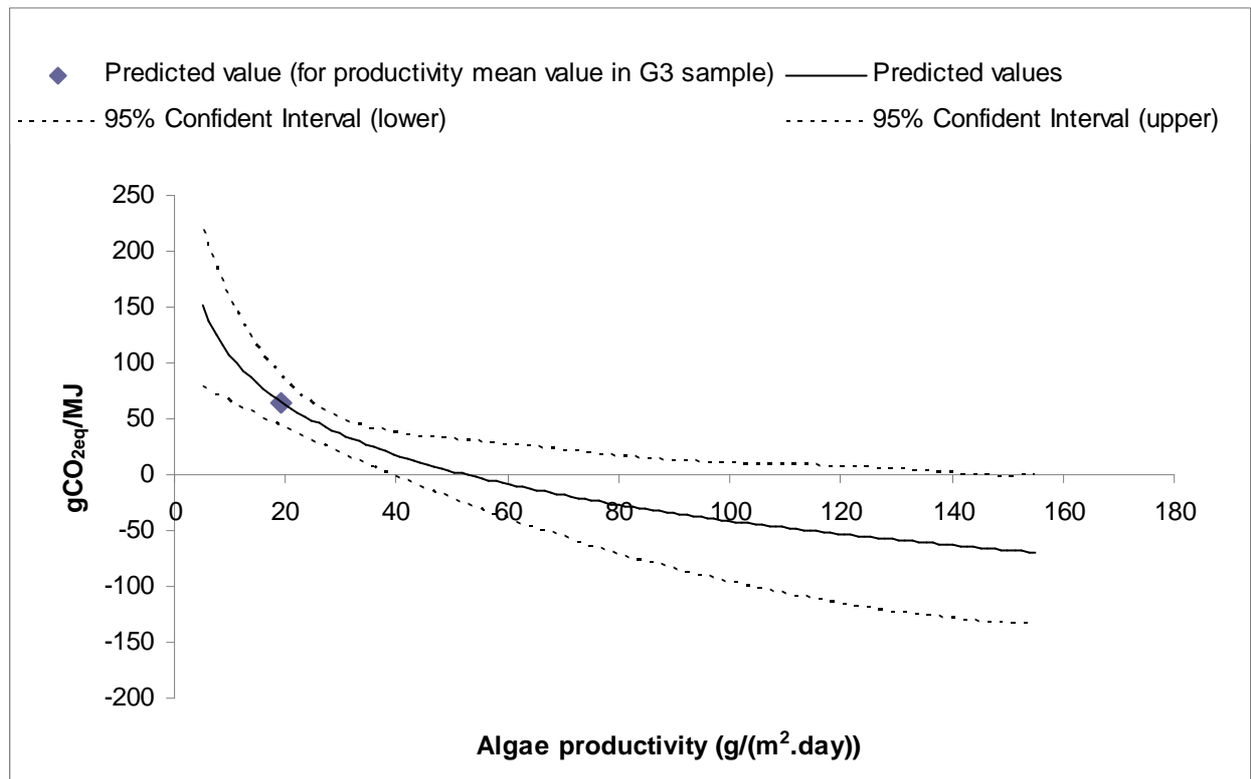


Figure 9 – Influence of algae productivity on predicted values of the  $e$ -s for G3 sample (1aG3 meta-model)

The same type of sensitivity analysis is conducted to test the influence of mass yield values of conversion process unit on BtL and Ethanol GHG emission values. As depicted in Figure 10 and Figure 11, mass yield value has a non linear effect on LCA GHG emissions of G2 biofuels, *ceteris paribus*. Variations for high values have less effect on the  $e$ -s than variations for low values. In addition, CIs are smaller for mass yield values around mean values than for extreme values, as previously described in the G3 sample.

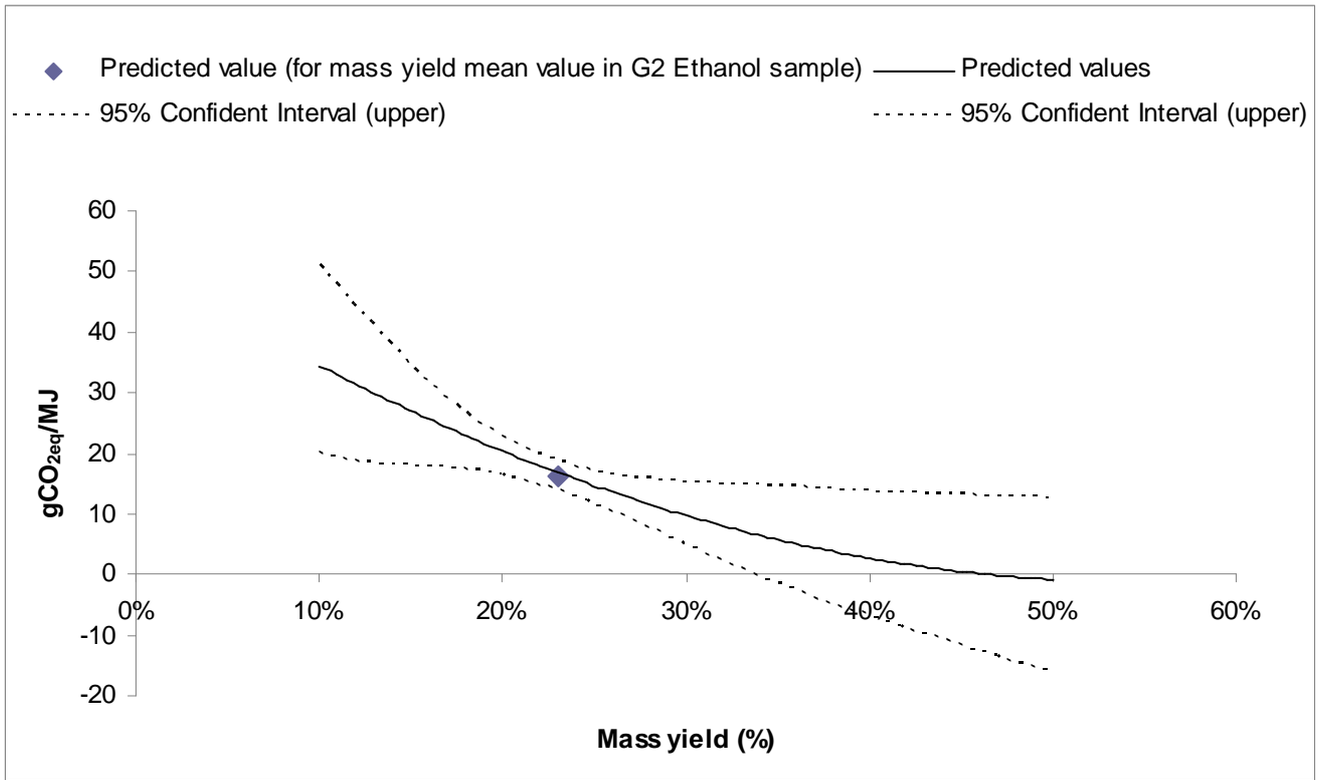


Figure 10 - Influence of the mass yield on predicted values of the  $e$ -s for Ethanol sample (1aEtha meta-model)

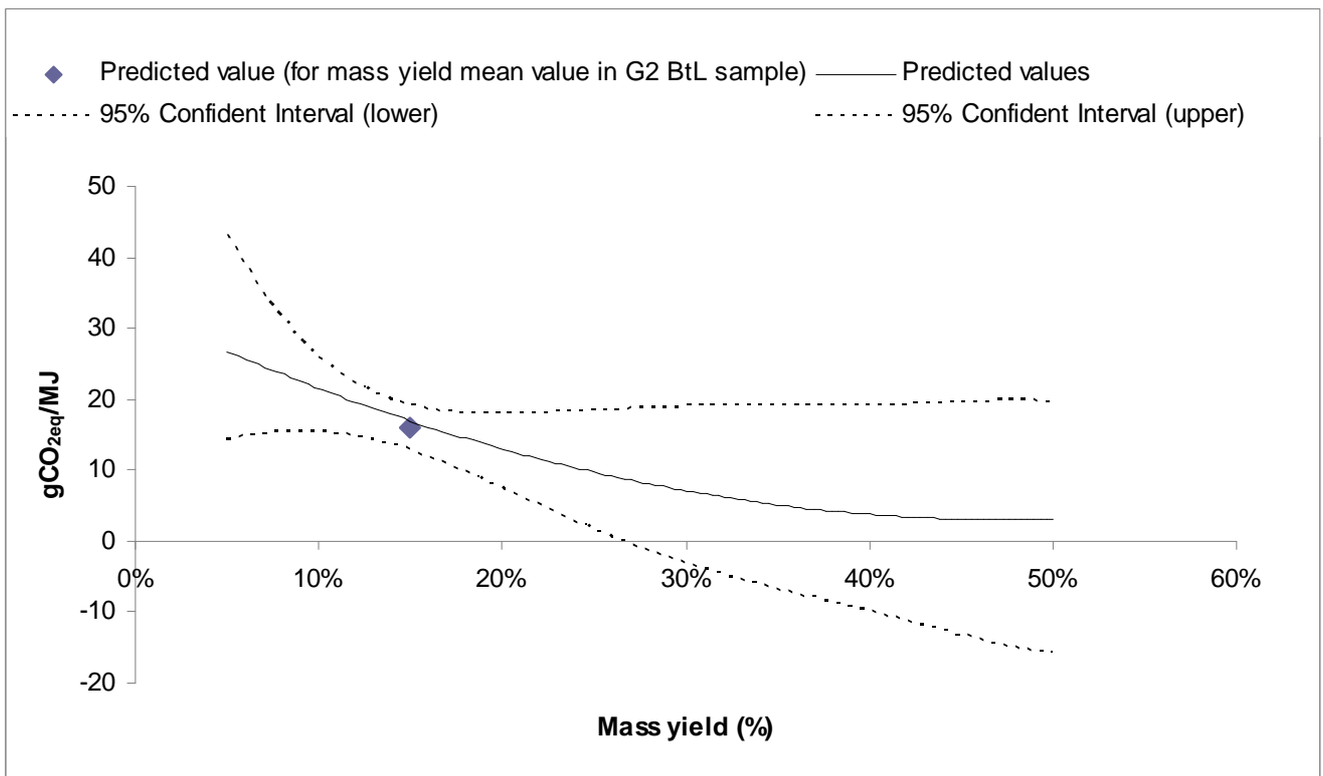


Figure 11 - Influence of the mass yield on predicted values of the  $e$ -s for BtL sample (1aBtL meta-model)

## 5 Concluding remarks and discussion

This article aims at synthesizing the literature of LCA studies which have estimated GHG emissions of advanced biofuels. Our literature review showed a high variation among the results (Figure 1). Thus, one can wonder *i*) if there is a consensus about GHG emission benefits from advanced biofuels and *ii*) why there is so much variation among results. To do so, we have been chosen to apply a specific *MA* methodology (the "*meta-regression analysis*", *MRA*) rather than a more classical narrative literature review approach. It provides a multivariate statistical analysis of previous estimated results to synthesize the available information. This assessment brings an extensive overview and contributes for a better understanding of the main factors inducing GHG emission variations. By using this original quantitative research framework, this article attempts to take the analysis of advanced biofuel GHG emissions one step further by complementing the qualitative surveys which have already been published [8–13]. We investigate – through an application – the potential for *MRA* to synthesize LCA literature by highlighting the main determinants of result variability in order to perform harmonization.

Our primary purpose was to identify and quantify which factors among *i*) technical data/characteristics, *ii*) author's methodological choices and *iii*) typology of the study under consideration have an impact on variations of the GHG emission estimates. Our results indicate a hierarchy between G3 and G2 biofuels: GHG emissions of G3 biofuels are statistically higher than those of Ethanol which, in turn, are superior to those of BtL. Moreover, whatever the type of advanced biofuel considered, North-American estimates are statistically higher than European estimates. Regarding author methodological choices, we have shown that some variables can influence the LCA results, such as the type of LCA approach (A-LCA vs. C-LCA), the method to account for coproducts and the fact of taking into account iLUC. Some technical variables appear to have an influence on GHG emission estimates. Concerning G2 biofuels, the mass yield has a negative and non-linear effect for both Ethanol and BtL whereas the type of process has a statistically significant effect only for BtL. For G3 biofuels, the algae productivity and its oil content have systematically a negative and non-linear effect. Finally, conclusions can be drawn also for some variables that have not been identified as variables influencing the final LCA result such as the type of biomass pretreatment in the Ethanol conversion process and the use of CCS in the BtL conversion process. The former is probably not statistically significant because most of the Ethanol technical data used in the different studies are derived from one single study [92]. The latter is a variable expected to have a negative impact in the GHG emission results but that could not be tested because all observations with the use of CCS fell in the outliers category.

The secondary purpose of this study was to address the harmonization issue in the field of advanced biofuel GHG emissions by using the technique of *benefits transfer using meta-regression models*. Our results may be summarized as follows. For each type of biofuel, a mean value of life cycle GHG emissions (expressed in g CO<sub>2</sub>eq / MJ of biofuel) weighted by the influence of its main drivers and its corresponding Confidence Interval is provided (Figure 5): about 60.0 (ranging from 43.3 to 76.7) for G3 biofuels; 19.7 (ranging from 17.4 to 22.0) for Ethanol; and 19.5 (ranging from 16.7 to 22.2) for BtL. Lastly, these values appear systematically higher for North-American estimates compared to those of Europe, *ceteris paribus* (Figure 7). Note that this range of values is lower than the fossil reference (about 83.8 in g CO<sub>2</sub>eq / MJ). However, only Ethanol and BtL do comply with the GHG emission reduction thresholds defined in both the US and EU directives.

Some results highlighted in this *MRA* have revealed some new information not previously assessed in this literature such as the existence of some non-linear effects regarding technical variables. Moreover, *MRA* provide *i*) a measure of the mean *e-s* and *ii*) a measure of the precision of this mean value estimate as provided by the corresponding Confidence Intervals.

Systematic reviews of LCA studies have gained interest due to their potential to clarify the impacts of particular products or services, producing more robust and policy-relevant results [14]. Most of the published so-called LCA meta-analyses rely on a "harmonization" procedure adjusting other study estimates based on "more consistent methods and assumptions" [15]. These studies typically harmonize technical parameters and methodological choices such as system boundaries, allocation procedures, impact calculation

method, etc. [17,33,82,86,97–100]. All the cited studies are able to reduce variability in calculated outcomes representing a useful starting point for more precise estimates of LCA results. However, this does not mean that this "harmonization" procedure, which we will call "normalization", produces more accurate results since the "more consistent methods and assumptions" applied are subjective depending on an author choices (different authors can consider different methods and assumptions to be more consistent).

The *MA* approach applied in this study is quite different and follows the traditional practice in biomedical sciences or economics, based on in-depth application of statistical methods and analysis. To our knowledge, Bureau *et al.* [34] are the only authors to use this type of approach in LCA systematic reviews. In the same way as their study, our results show that this methodology can be consistently applied for the identification of parameters that influence a biofuel LCA result. Moreover, we have gone further by proposing a method to predict LCA results using a meta-model. This can be seen as a harmonization method alternative to the one applied currently in LCA *meta-analysis* ("normalization").

Our meta-model is obtained from a meta-regression and, therefore, it contains the parameters that were statistically proven to influence LCA results in a given sample. Our results show that, with this approach, we can provide more than a mean value and an interquartile range for the *e-s* as done in the "normalization" studies. We can calculate a real confidence interval for our predictions.

Among the LCA *meta-analysis* reviewed, the only meta-model built using regression techniques, is attempted in a work from Padey *et al.* [101]. However, their wind turbine LCA meta-model is based on a limited number of parameters (lifetime and wind speed) and their regression method cannot account for qualitative parameters (dummy variables used in this study). Therefore, we observe that this is a field where significant progress can be made and we would recommend that the LCA community should work more closely with the Econometrics community so that more studies of this type could be conducted.

However, there are many limitations typically associated with *MA*. In the construction of the database, for example, there is always some exogenous information that has to be provided. Even if we avoid it as much as possible, in some cases it is necessary. This happened especially in the calculation of the *e-s* where the data required for the conversion of units (LHV, density, motor performance, etc.) was not always provided by the study in question.

Moreover, there is a compromise that has to be made between the number of studies that pass the screening process and the number of independent variables that are used in the description of an observation. In a *MA* database, all of the observations in a given sample have to be described with the same amount of independent variables. Theoretically, all the parameters that potentially influence the *e-s* have to be included. However, in LCA, the results are affected by hundreds of inputs and methodological choices, making it impossible to fully explain all the results of a big number of observations given the heterogeneity in LCA reporting. It was our judgement and experience in conducting LCA studies, but also previous narrative surveys, that determined which explanatory variables should be included in the database.

Finally, there may be some limitations regarding the statistical population of the *MA* sample. Heath and Mann [15] highlight the fact that *MA* cannot make up for a lack of studies on a certain technology or methodological issue. In our case, for example, there are only 3 observations for BtL including CCS in its production pathway and these were coincidentally discarded from the meta-regression sample as outliers. Therefore, no conclusions could be drawn from this technological parameter. Another example is the limited number of consequential LCAs, also limiting the conclusions we can reach concerning this methodological choice.

On our view, *MA* appear thus more as a complementary methodology than an alternative one to more classical narrative surveys.

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# Appendices

## Appendix A. Technical description of advanced biofuels

Figure A.1 represents the main steps involved in the production of second and third generation biofuels (G2 and G3 biofuels respectively) discussed in this paper and the following text contain a brief description of their production processes.

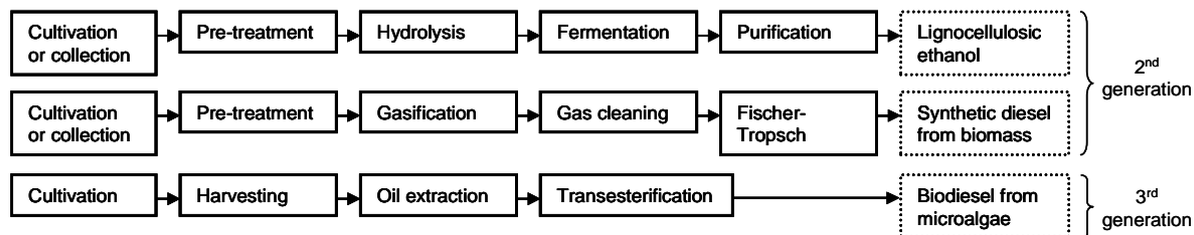


Figure A.1 – Main steps in the production of advanced biofuels

Second generation Ethanol is obtained from the biochemical conversion of annual crop residues (e.g. corn stover) and perennial crops (e.g. miscanthus). A pretreatment of the biomass is necessary to separate the cellulose from hemicellulose and lignin. Once the cellulose is accessible, enzymes are used to hydrolyze these molecules, transforming them into sugars that can be fermented. The product of fermentation needs to be distilled and dehydrated for obtaining pure Ethanol [92,102].

Synthetic diesel from biomass is also known as BtL (Biomass to Liquids) or biomass FT-diesel. It is produced by the thermochemical conversion of forest residues, herbaceous energy crops (e.g. switchgrass) and woody biomass (e.g. poplar), which is a promising second generation pathway. A pretreatment of the biomass is necessary so that it can be loaded into the gasifier. In the gasifier, the biomass suffers a thermal treatment into what is known as "syngas", composed mainly of H<sub>2</sub> and CO. Impurities are removed from the "syngas" during a gas cleaning step, due to the high sensibility of the Fischer-Tropsch (FT) reaction catalyst. The synthetic diesel is obtained after the upgrading (hydrocracking) of the products from the FT unit [103,104].

Biodiesel can be produced from conventional transesterification of oil extracted from microalgae that have a higher theoretical productivity per hectare than conventional vegetable oil crops (e.g. soybeans, palm). Microalgae can be cultivated in open ponds or photobioreactors (PBR) and the technologies for harvesting, drying and extracting oil still require considerable research effort. Various pathways are studied in order to reduce costs and energy consumption in the production process. The use of power plant flue gas as a CO<sub>2</sub> source for growing algae or wastewater as a source of nutrients are potential options for this biodiesel pathway [95,105].

Studies about hydrotreated vegetable oil (HVO) from the hydrogenation of microalgae oil were also included in this literature review. It has different characteristics than biodiesel but the most important life cycle steps involving microalgae growth, harvesting and oil extraction are the same. HVO, as well as BtL, are being studied as renewable alternatives not just for road transportation but also for the aviation industry.

## Appendix B. Complements on the database

**Table B.1 - List of selected LCA studies assessing GHG emissions of G2 and G3 biofuels**

Name	Authors	Title	Year
Bai et al. (2010)	Bai Y, Luo L, van der Voet E	Life cycle assessment of switchgrass derived ethanol as transport fuel	2010
Batan et al. (2010)	Batan L, Quinn J, Willson B, Bradley T	Net energy and greenhouse emission evaluation of biodiesel derived from microalgae	2010
Campbell et al. (2010)	Campbell PK, Beer T, Batten D	Life cycle assessment of biodiesel production from microalgae in ponds (submitted)	2010
Cherubini et al. (2011)	Cherubini F, Hammer Stromman A, Ulgiati S	Influence of allocation methods on the environmental performance of biorefinery products - A case study	2011
Choudhury et al. (2002)	Choudhury R, Weber T, Schindler J, Weindorf W, Wurster R	Well-to-wheel analysis of energy use and greenhouse gas emissions of advanced fuel/vehicle systems	2002
Delucchi (2006)	Delucchi M	Life Cycle Analysis of Biofuels	2006
Dussault et al. (2010)	Dussault P.C, Bradt L, Ponce-Ortega J.M, El-Halwagi M.M	Incorporation of process integration into life cycle analysis for the production of biofuels	2010
Elsayed et al. (2003)	Elsayed M.A, Matthews R, Mortimer N.D	Carbon and Energy Balances for a Range of Biofuels Options	2003
Fazio & Monti (2011)	Fazio S, Monti A	Life cycle assessment of different bioenergy production systems including perennial and annual crops	2011
Gonzales-Garcia et al. (2009a)	Gonzales Garcia S, Luo L, Moreira T, Feijoo G, Huppes G	Life cycle assessment of flax shives derived second generation ethanol fueled automobiles in Spain (submitted)	2009
Gonzales-Garcia et al. (2009b)	Gonzales-Garcia S, Gasol C.M, Gabarrelle X, Rieradevall J, Moreira T, Feijoo G	Environmental profile of ethanol from poplar biomass as transport fuel in southern Europe (submitted)	2009
Gonzalez-Garcia et al. (2009c)	Gonzalez-Garcia S, Moreira M.T, Feijoo G	Environmental performance of lignocellulosic bioethanol production from Alfalfa stems(submitted)	2009
Groode et al. (2007)	Groode T.A, Heywood J.B	Ethanol a look ahead	2007
Haase et al. (2009)	Haase M, Skott S, Fröhling M	Ecological evaluation of selected 1st and 2nd generation biofuels- FT fuel from wood and ethanol from sugar beets (submitted)	2009
Hoefnagels et al. (2010)	Hoefnagels R, Smeets E, Faaij A	Greenhouse gas footprints of different biofuel production systems	2010
Hsu et al. (2010)	Hsu DD, Inman D, Heath G, Wolfrum E, Mann MK, Aden A	Life Cycle Environmental Impacts of Selected U.S. Ethanol Production and Use Pathways in 2022	2010
JEC (2007)	JEC - Joint Research Centre-EUCAR-CONCAWE collaboration	Well to Wheels analysis of future automotive fuels and powertrains in the European Context (version2c)	2007
JEC (2011)	JEC - Joint Research Centre-EUCAR-CONCAWE collaboration	Well to Wheels analysis of future automotive fuels and powertrains in the European Context (version3c)	2011
Jungbluth et al. (2007)	Jungbluth N, Frischknecht R, Emmenegger MF, Steiner R, Tuschshmid M	Life cycle assessment of BTL-fuel production: life cycle impact assessment and interpretation. Renewable Fuels for Advanced Powertrains (RENEW) Project	2007
Jungbluth et al. (2008)	Jungbluth N, Busser S, Frischknecht R, Tuschschmid M	Life cycle assessment of biomass-to-liquid fuels	2008
Kaufman et al. (2010)	Kaufman A.S, Meier PJ, Sinistore JC, Reinemann DJ	Applying life -cycle assessment to low carbon fuel standards -How allocation choices influence carbon intensity for renewable transportation fuels	2010
Koponen et al. (2009)	Koponen K, Soimakallio S, Sipilä E	Assessing greenhouse gas emissions of waste-derived ethanol in accordance with the EU RED methodology for biofuels	2009
Lardon et al. (2009)	Lardon L, Helias A, Sialve B, Steyer JP, Bernard O	Life cycle assessment of biodiesel from microalgae	2009
Luo et al. (2009)	Luo L, van der Voet E, Huppes G	Allocation issues in LCA methodology : a case study of corn-stover based fuel ethanol	2009
McKechnie et al. (2011)	McKechnie J, Zhang Y, Ogino A, Saville B, Sleep S, Turner M, Pontius R, MacLean H.L	Impact of co-location, co-production, and process energy source on life cycle energy use and greenhouse gas emissions of lignocellulosic ethanol	2011
Mehlin et al. (2003)	Mehlin M, Zauner M, Gühnemann A, Aoki R, Vance C	Renewable Fuels for Cross Border Transportation	2003
Mu et al. (2010)	Mu D, Seager T, Rao P.S	Comparative life cycle assessment of lignocellulosic ethanol production : Biochemical versus thermochemical conversion	2010
Mullins et al. (2010)	Mullins K.A, Griffin W.M, Matthews H.S	Policy implications of uncertainty in modeled life cycle greenhouse gas emissions of biofuels (submitted)	2010
RED (2009)	European Parliament	Directive 2009/28/EC of the European Parliament and of the Council of 23 April 2009 On the promotion of the use of energy from renewable sources	2009
RFS2 (2010)	United States Environmental Protection Agency	Renewable fuel standard program (RFS2) regulatory impact analysis	2010
Sander et al. (2010)	Sander K, Murthy GS	Life cycle analysis of algae biodiesel	2010
Schmitt et al. (2011)	Schmitt E, Bura R, Gustafson R, Cooper J, Vajzovic A	Converting lignocellulosic solid waste into ethanol for the State of Washington: An investigation of treatment technologies and environmental impacts (submitted)	2011
Sheehan et al. (2004)	Sheehan J, Aden A, Paustian K, Killian K, Brenner J, Walsh M, Nelson R	Energy and environmental aspects of using corn stover for fuel ethanol	2004
Spatari et al. (2005)	Spatari S, Zhang Y, MacLean H.L	Life cycle assesment of switchgrass and corn-stover derived ethanol-fueled automobiles	2005
Spatari et al. (2009)	Spatari S, Bagley DM, MacLean HL	Life cycle evaluation of emerging lignocellulosic ethanol conversion technologies (submitted)	2009
Spatari et al. (2010)	Spatari S, MacLean H.L	Characterizing models uncertainties in the life cycle of lignocellulose-based ethanol fuels	2010
Stephenson et al. (2010a)	Stephenson A.L, Dupree P, Scott S.A, Dennis J.S	The environmental and economic sustainability of potential bioethanol from willow in the UK	2010
Stephenson et al. (2010b)	Stephenson A.L, Kazamia E, Dennis J.S, Howe C.J, Scott S.A, Smith A.G	Life cycle assesment of potential algal biodiesel production in the United Kingdom : A comparison of raceways and air-lift tubular bioreactors	2010
Stichnothe et al. (2009)	Stichnothe H, Azapagic A	Bioethanol from waste: life cycle estimation of the greenhouse gas saving potential	2009
Stratton et al. (2010)	Stratton RW, Wong HM, Hilman JI	Life Cycle Greenhouse Gas emissions from alternative jet fuels	2010
van Vliet et al. (2009)	van Vliet O.P.R, Faaij A.P.C, Turkenburg W.C	Fischer-Tropsch diesel production in a well-to wheel perspective : a carbon, energy flow and cost analysis	2009
Vera-Morales et al. (2009)	Vera-Morales M, Schäfer A	Well-to-wheels analysis of future automotive fuels and powertrains in the european context	2009
Wang et al. (2010)	Wang M, Huo H, Arora S	Methods of dealing with co-products of biofuels in life cycle analysis and consequent results within the U.S context (submitted)	2010
Wang et al. (2011)	Wang M.Q, Han J, Haq Z, Tyner W.E, Wu M, Elgowainy A	Energy and greenhouse gas emission effects of corn and cellulosic ethanol with technology improvements and land use changes	2011
Whittaker et al. (2011)	Whittaker C, McManus M.C, Hammond G.P	Greenhouse gas reporting for biofuels : a comparison between RED, RFTO and PAS2050 methodologies	2011
Wu et al. (2005)	Wu M, Wu Y, Wang M	Mobility Chains Analysis of Technologies for Passenger Cars and Light-Duty Vehicles Fueled with Biofuels: Application of the GREET Model to the Role of Biomass in America's Energy Future (RBAEF) Project	2005
Xie et al. (2011)	Xie X, Wang M, Han J	Assessment of Fuel-Cycle Energy Use and Greenhouse Gas Emissions for Fischer-Tropsch Diesel from Coal and Cellulosic Biomass	2011

**Table B.2 - Description of variables included in the database**

Variables Family	Description of variables	Variables	Type of variable	Unit
<b>Technical data</b>				
Type of biofuel	Second generation biofuel	<i>gen_2</i>	Binary (=1 if 'true'; 0 else)	
	Third generation biofuel	<i>gen_3</i>	Binary (=1 if 'true'; 0 else)	
	Ethanol	<i>etha</i>	Binary (=1 if 'true'; 0 else)	
	Biomass to Liquid (BtL)	<i>btl</i>	Binary (=1 if 'true'; 0 else)	
	Fatty Acid Methyl Ester (FAME)	<i>fame</i>	Binary (=1 if 'true'; 0 else)	
	Hydrotreated Vegetable Oil (HVO)	<i>hvo</i>	Binary (=1 if 'true'; 0 else)	
Type of biomass feedstock	Microalgae	<i>mat_algae</i>	Binary (=1 if 'true'; 0 else)	
	Agricultural residues	<i>mat_agrires</i>	Binary (=1 if 'true'; 0 else)	
	Forestry residues	<i>mat_for</i>	Binary (=1 if 'true'; 0 else)	
	Energy crop	<i>mat_energcult</i>	Binary (=1 if 'true'; 0 else)	
	Farmed wood	<i>mat_farmwood</i>	Binary (=1 if 'true'; 0 else)	
	Others	<i>mat_other</i>	Binary (=1 if 'true'; 0 else)	
	Cultivated feedstock	<i>mat_cult</i>	Binary (=1 if 'Cultivated feedstock'; 0 if 'Waste feedstock')	
Type of coproducts	Existence of coproduct	<i>coprod</i>	Binary (=1 if 'true'; 0 else)	
	Glycerin as coproduct	<i>cop_gly</i>	Binary (=1 if 'true'; 0 else)	
	Electricity as coproduct	<i>cop_elec</i>	Binary (=1 if 'true'; 0 else)	
	Heat as coproduct	<i>cop_heat</i>	Binary (=1 if 'true'; 0 else)	
	Algal meal as coproduct	<i>cop_algmeal</i>	Binary (=1 if 'true'; 0 else)	
	Biogas as coproduct	<i>cop_biog</i>	Binary (=1 if 'true'; 0 else)	
	Other coproduct	<i>cop_other</i>	Binary (=1 if 'true'; 0 else)	
	Other coproduct than electricity	<i>cop_otherelec</i>	Binary (=1 if 'true'; 0 else)	
	Number of coproducts	<i>coprod_num</i>	Binary (=1 if 'true'; 0 else)	
Type of technologies and associated yields	Mass yield provided	<i>mass_yield_exist</i>	Binary (=1 if 'true'; 0 else)	
	Value of mass yield	<i>g2_mass_yield</i>	Quantitative	% mass
	Steam explosion as biomass pretreatment for Ethanol technology	<i>eth_expl</i>	Binary (=1 if 'true'; 0 else)	
	Dilute sulfuric acid as biomass pretreatment for Ethanol technology	<i>eth_ac</i>	Binary (=1 if 'true'; 0 else)	
	Ammonia fibre explosion as biomass pretreatment for Ethanol technology	<i>eth_amm</i>	Binary (=1 if 'true'; 0 else)	
	Other processes as biomass pretreatment for Ethanol technology	<i>eth_other</i>	Binary (=1 if 'true'; 0 else)	
	Torrefaction as biomass pretreatment for BtL technology	<i>btl_pre_torr</i>	Binary (=1 if 'true'; 0 else)	
	Pyrolyse as biomass pretreatment for BtL technology	<i>btl_pre_pyro</i>	Binary (=1 if 'true'; 0 else)	
	No biomass pretreatment for BtL technology	<i>btl_pre_none</i>	Binary (=1 if 'true'; 0 else)	
	Autothermic BtL technology	<i>btl_pro_autoth</i>	Binary (=1 if 'true'; 0 else)	
	Allothermic BtL technology with natural gas as fuel	<i>btl_pro_alng</i>	Binary (=1 if 'true'; 0 else)	
	Allothermic BtL technology with imported electricity as fuel	<i>btl_pro_alelec</i>	Binary (=1 if 'true'; 0 else)	
	Allothermic BtL technology with renewable energy as fuel	<i>btl_pro_alrenew</i>	Binary (=1 if 'true'; 0 else)	
	BtL technology with tail gas recycled	<i>btl_gasrecycl</i>	Binary (=1 if 'true'; 0 else)	
	BtL technology with Carbon Capture and Storage	<i>btl_ccs</i>	Binary (=1 if 'true'; 0 else)	
	Microalgae productivity for G3 biofuel	<i>g3_productivity</i>	Quantitative	g/(m <sup>2</sup> .day)
	Oil content of microalgae for G3 biofuel	<i>g3_oil</i>	Quantitative	% dry mass
	Open Pond for microalgae cultivation for G3 biofuel	<i>g3_Oppond</i>	Binary (=1 if 'open pond'; 0 if "photobioreactor or hybrid")	
	Geographical location of the case study	North America	<i>zloc_us</i>	Binary (=1 if 'true'; 0 else)
Europe		<i>zloc_eu</i>	Binary (=1 if 'true'; 0 else)	
Other		<i>zloc_other</i>	Binary (=1 if 'true'; 0 else)	

Methodological choices				
Type of LCA approach	Attributional LCA	<i>lca_att</i>	Binary (=1 if 'true'; 0 else)	
	Consequential LCA	<i>lca_cons</i>	Binary (=1 if 'true'; 0 else)	
System boundaries	Well To Tank	<i>wtt</i>	Binary (=1 if 'true'; 0 else)	
	Well To Wheel	<i>wtw</i>	Binary (=1 if 'true'; 0 else)	
	Infrastructures taken into account in boundaries	<i>infrastruct</i>	Binary (=1 if 'true'; 0 else)	
Method for taking into account coproducts	Energetic allocation	<i>copval_alloc_ener</i>	Binary (=1 if 'true'; 0 else)	
	Mass allocation	<i>copval_alloc_mass</i>	Binary (=1 if 'true'; 0 else)	
	Economic allocation	<i>copval_alloc_markval</i>	Binary (=1 if 'true'; 0 else)	
	Energetic allocation	<i>copval_alloc_exerg</i>	Binary (=1 if 'true'; 0 else)	
	Allocation	<i>copval_alloc</i>	Binary (=1 if 'true'; 0 else)	
	System expansion	<i>copval_systexp</i>	Binary (=1 if 'true'; 0 else)	
	Hybrid	<i>copval_hyb</i>	Binary (=1 if 'true'; 0 else)	
Carbon neutral	Carbon neutral hypothesis for WTW G3 biofuel studies	<i>wtw*g3_carbneut</i>	Binary (=1 if 'true'; 0 else)	
Characterization method for impact assessment	Number of greenhouse gases taken into account	<i>gas_num</i>	Binary (=1 if 'number >3'; 0 else)	
Method for taking into account N2O emission from N input	Use of IPCC method	<i>ass_ipcc</i>	Binary (=1 if 'true'; 0 else)	
Method for taking into account Land Use Change	LUC taken into account	<i>luc</i>	Binary (=1 if 'true'; 0 else)	
	Direct LUC taken into account	<i>luc_dir</i>	Binary (=1 if 'true'; 0 else)	
	Indirect LUC taken into account	<i>luc_indir</i>	Binary (=1 if 'true'; 0 else)	
Method for taking into account uncertainties	No method for taking into account uncertainties	<i>uncer_ref</i>	Binary (=1 if 'true'; 0 else)	
	Use of Monte Carlo analysis	<i>uncer_MC</i>	Binary (=1 if 'true'; 0 else)	
	Use of sensitivity analysis	<i>uncer_SA</i>	Binary (=1 if 'true'; 0 else)	
	Use of a method for taking into account uncertainties	<i>uncer_MCSA</i>	Binary (=1 if 'true'; 0 else)	
Number and type of environmental impact indicator assessed in the study	Net Energy Value assessed	<i>impcat_nev</i>	Binary (=1 if 'true'; 0 else)	
	Non Renewable Energy Consumption assessed	<i>impcat_nrc</i>	Binary (=1 if 'true'; 0 else)	
	Energetic indicator assessed	<i>impcat_nrcnev</i>	Binary (=1 if 'true'; 0 else)	
	Global Warming indicator assessed	<i>impcat_gwp</i>	Binary (=1 if 'true'; 0 else)	
	Other environmental indicator assessed	<i>impcat_other</i>	Binary (=1 if 'true'; 0 else)	
	Number of environmental indicator assessed	<i>impcat_all</i>	Quantitative	number
Typology of the study				
Type of study	Peer review literature	<i>lit_pr</i>	Binary (=1 if 'true'; 0 else)	
	Official report	<i>lit_or</i>	Binary (=1 if 'true'; 0 else)	
	Directive or Standard	<i>lit_dir</i>	Binary (=1 if 'true'; 0 else)	
	Official report or directive/standard	<i>lit_ordir</i>	Binary (=1 if 'true'; 0 else)	
	Working paper	<i>lit_wp</i>	Binary (=1 if 'true'; 0 else)	
Year of publication	Year of publication	<i>year</i>	Quantitative	year
	Year of publication after 2007	<i>year_07</i>	Binary (=1 if 'true'; 0 else)	
	Year of publication after 2009	<i>year_09</i>	Binary (=1 if 'true'; 0 else)	
	Year of publication after 2010	<i>year_10</i>	Binary (=1 if 'true'; 0 else)	
Geographical location of authors	North America	<i>zlab_us</i>	Binary (=1 if 'true'; 0 else)	
	Europe	<i>zlab_eu</i>	Binary (=1 if 'true'; 0 else)	
	Other	<i>zlab_other</i>	Binary (=1 if 'true'; 0 else)	

**Table B.3 - Share of studies and observations for the geographical location of the authors and of assessed pathways**

Weighted by	Geographical location of the authors			Geographical location of assessed pathways		
	North America	Europe	Other	North America	Europe	Other
Studies (47)	45%	53%	2%	-	-	-
Observations (593)	32%	67%	1%	34%	64%	2%

**Table B.4 – Descriptive statistics for the effect size and variables of the original Whole sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	593	34.45	89.34	-142.18	1377.90
es_min	593	-11.25	70.81	-352.81	881.44
es_max	593	80.18	155.52	-127.01	1874.37
es_var	593	2264.65	9868.76	0.00	64159.87
es_et	593	23.37	41.49	0.03	253.30
<b>Technical data</b>					
gen_2	593	0.87	0.34	0	1
gen_3	593	0.13	0.34	0	1
etha	593	0.61	0.49	0	1
btl	593	0.26	0.44	0	1
fame	593	0.11	0.31	0	1
hvo	593	0.03	0.16	0	1
mat_algae	593	0.13	0.34	0	1
mat_agrires	593	0.22	0.41	0	1
mat_for	593	0.05	0.22	0	1
mat_enercult	593	0.28	0.45	0	1
mat_farmwood	593	0.10	0.30	0	1
mat_other	593	0.23	0.42	0	1
mat_cult	593	0.38	0.49	0	1
coprod	593	0.91	0.29	0	1
cop_gly	593	0.09	0.29	0	1
cop_elec	593	0.63	0.48	0	1
cop_heat	593	0.12	0.32	0	1
cop_algmeal	593	0.10	0.30	0	1
cop_biog	593	0.18	0.39	0	1
cop_other	593	0.25	0.54	0	2
cop_otherelec	593	0.48	0.50	0	1
coprod_num	593	1.36	1.00	0	5
mass_yield_exist	593	0.64	0.48	0	1
g2_mass_yield	378	0.22	0.07	0.07	0.50
eth_expl	361	0.08	0.28	0	1
eth_ac	361	0.83	0.37	0	1
eth_amm	361	0.08	0.28	0	1
eth_other	361	0.00	0.00	0	0
btl_pre_torr	153	0.07	0.25	0	1
btl_pre_pyro	153	0.05	0.21	0	1
btl_pre_none	153	0.89	0.32	0	1
btl_pro_autoth	153	0.86	0.35	0	1
btl_pro_alng	153	0.08	0.27	0	1
btl_pro_alelec	153	0.03	0.18	0	1
btl_pro_alrenew	153	0.03	0.16	0	1
btl_gasrecycl	153	0.01	0.11	0	1
btl_ccs	153	0.02	0.14	0	1
g3_productivity	76	30.09	18.86	6.85	150.00
g3_oil	77	0.41	0.11	0.15	0.70
g3_Oppond	77	0.61	0.49	0	1
zloc_us	593	0.34	0.47	0	1
zloc_eu	593	0.64	0.48	0	1
zloc_other	593	0.02	0.15	0	1
<b>Methodological choices</b>					
lca_att	593	0.97	0.17	0	1
lca_cons	593	0.03	0.17	0	1
wtt	593	0.38	0.49	0	1
wtw	593	0.62	0.49	0	1
infrastruct	593	0.44	0.50	0	1
copval_alloc_ener	593	0.26	0.44	0	1
copval_alloc_mass	593	0.04	0.20	0	1
copval_alloc_markval	593	0.12	0.32	0	1
copval_alloc_exerg	593	0.01	0.07	0	1
copval_alloc	593	0.42	0.49	0	1
copval_systexp	593	0.55	0.50	0	1
copval_hyb	593	0.07	0.26	0	1
wtw*g3_carbneut	77	0.00	0.00	0	0
gas_num	394	0.26	0.44	0	1
ass_ipcc	573	0.37	0.48	0	1
luc	593	0.51	0.50	0	1
luc_dir	593	0.51	0.50	0	1
luc_indir	593	0.03	0.18	0	1
uncer_ref	593	0.57	0.50	0	1
uncer_MC	593	0.10	0.30	0	1
uncer_SA	593	0.38	0.49	0	1
uncer_MCSEA	593	0.48	0.50	0	1
impcat_nev	593	0.31	0.46	0	1
impcat_nrc	593	0.41	0.49	0	1
impcat_nrcnev	593	0.50	0.50	0	1
impcat_gwp	593	1.00	0.00	1	1
impcat_other	593	1.09	2.28	0	9
impcat_all	593	2.81	2.43	1	12
<b>Typology of the study</b>					
lit_pr	593	0.63	0.48	0	1
lit_or	593	0.09	0.29	0	1
lit_dir	593	0.03	0.17	0	1
lit_ordir	593	0.12	0.32	0	1
lit_wp	593	0.25	0.44	0	1
year	593	2009.38	1.31	2002	2011
year_07	593	0.97	0.17	0	1
year_09	593	0.90	0.30	0	1
year_10	593	0.56	0.50	0	1
zlab_us	593	0.32	0.47	0	1
zlab_eu	593	0.67	0.47	0	1
zlab_other	593	0.01	0.10	0	1

**Table B.5 – Descriptive statistics for the effect size and variables of the original G3 sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	77	88.87	211.85	-96.47	1377.90
es_min	77	-46.68	146.41	-352.81	881.44
es_max	77	224.42	351.73	-7.55	1874.37
es_var	77	11898.53	23739.99	265.09	64159.87
es_et	77	69.16	84.91	16.28	253.30
<b>Technical data</b>					
gen_2	77	0.00	0.00	0	0
gen_3	77	1.00	0.00	1	1
etha	77	0.00	0.00	0	0
btl	77	0.00	0.00	0	0
fame	77	0.82	0.39	0	1
hvo	77	0.19	0.40	0	1
mat_algae	77	1.00	0.00	1	1
mat_agrires	77	0.00	0.00	0	0
mat_for	77	0.00	0.00	0	0
mat_enercult	77	0.00	0.00	0	0
mat_farmwood	77	0.00	0.00	0	0
mat_other	77	0.00	0.00	0	0
mat_cult	77	0.00	0.00	0	0
coprod	77	1.00	0.00	1	1
cop_gly	77	0.71	0.45	0	1
cop_elec	77	0.27	0.45	0	1
cop_heat	77	0.40	0.49	0	1
cop_algmeal	77	0.79	0.41	0	1
cop_biog	77	0.00	0.00	0	0
cop_other	77	0.95	0.93	0	2
cop_otherelec	77	0.87	0.34	0	1
coprod_num	77	2.99	1.71	1	5
mass_yield_exist	77	0.05	0.22	0	1
g3_productivity	76	30.09	18.86	6.85	150.00
g3_oil	77	0.41	0.11	0.15	0.70
g3_Oppond	77	0.61	0.49	0	1
zloc_us	77	0.42	0.50	0	1
zloc_eu	77	0.51	0.50	0	1
zloc_other	77	0.08	0.27	0	1
<b>Methodological choices</b>					
lca_att	77	0.92	0.27	0	1
lca_cons	77	0.08	0.27	0	1
wtt	77	0.65	0.48	0	1
wtw	77	0.35	0.48	0	1
infrastruct	77	0.51	0.50	0	1
copval_alloc_ener	77	0.08	0.27	0	1
copval_alloc_mass	77	0.00	0.00	0	0
copval_alloc_markval	77	0.03	0.16	0	1
copval_alloc_exerg	77	0.00	0.00	0	0
copval_alloc	77	0.10	0.31	0	1
copval_systexp	77	0.90	0.31	0	1
copval_hyb	77	0.55	0.50	0	1
wtw*g3_carbneut	77	0.00	0.00	0	0
gas_num	77	0.45	0.50	0	1
ass_ipcc	77	0.48	0.50	0	1
luc	77	0.08	0.27	0	1
luc_dir	77	0.08	0.27	0	1
luc_indir	77	0.08	0.27	0	1
uncer_ref	77	0.39	0.49	0	1
uncer_MC	77	0.00	0.00	0	0
uncer_SA	77	0.61	0.49	0	1
uncer_MCOSA	77	0.61	0.49	0	1
impcat_nev	77	0.30	0.46	0	1
impcat_nrc	77	0.40	0.49	0	1
impcat_nrcnev	77	0.70	0.46	0	1
impcat_gwp	77	1.00	0.00	1	1
impcat_other	77	0.66	1.13	0	5
impcat_all	77	2.36	1.38	1	7
<b>Typology of the study</b>					
lit_pr	77	0.73	0.45	0	1
lit_or	77	0.00	0.00	0	0
lit_dir	77	0.08	0.27	0	1
lit_ordir	77	0.08	0.27	0	1
lit_wp	77	0.19	0.40	0	1
year	77	2009.88	0.32	2009	2010
year_07	77	1.00	0.00	1	1
year_09	77	1.00	0.00	1	1
year_10	77	0.88	0.32	0	1
zlab_us	77	0.42	0.50	0	1
zlab_eu	77	0.51	0.50	0	1
zlab_other	77	0.08	0.27	0	1

**Table B.6 – Descriptive statistics for the effect size and variables of the original G2 sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	516	26.33	45.20	-142.18	518.40
es_min	516	-5.96	48.83	-307.59	195.45
es_max	516	58.65	77.27	-127.01	841.35
es_var	516	827.04	3582.78	0.00	27149.13
es_et	516	16.54	23.55	0.03	164.77
<b>Technical data</b>					
gen_2	516	1.00	0.00	1	1
gen_3	516	0.00	0.00	0	0
etha	516	0.70	0.46	0	1
btl	516	0.30	0.46	0	1
fame	516	0.00	0.00	0	0
hvo	516	0.00	0.00	0	0
mat_algae	516	0.00	0.00	0	0
mat_agrires	516	0.25	0.43	0	1
mat_for	516	0.06	0.23	0	1
mat_enercult	516	0.32	0.47	0	1
mat_farmwood	516	0.11	0.31	0	1
mat_other	516	0.26	0.44	0	1
mat_cult	516	0.43	0.50	0	1
coprod	516	0.89	0.31	0	1
cop_gly	516	0.00	0.00	0	0
cop_elec	516	0.69	0.46	0	1
cop_heat	516	0.08	0.27	0	1
cop_algmeal	516	0.00	0.00	0	0
cop_biog	516	0.21	0.41	0	1
cop_other	516	0.15	0.35	0	1
cop_otherelec	516	0.42	0.49	0	1
coprod_num	516	1.12	0.52	0	3
mass_yield_exist	516	0.73	0.44	0	1
g2_mass_yield	378	0.22	0.07	0.07	0.50
eth_expl	361	0.08	0.28	0	1
eth_ac	361	0.83	0.37	0	1
eth_amm	361	0.08	0.28	0	1
eth_other	361	0.00	0.00	0	0
btl_pre_torr	153	0.07	0.25	0	1
btl_pre_pyro	153	0.05	0.21	0	1
btl_pre_none	153	0.89	0.32	0	1
btl_pro_autoth	153	0.86	0.35	0	1
btl_pro_alng	153	0.08	0.27	0	1
btl_pro_alelec	153	0.03	0.18	0	1
btl_pro_alrenew	153	0.03	0.16	0	1
btl_gasrecycl	153	0.01	0.11	0	1
btl_ccs	153	0.02	0.14	0	1
zloc_us	516	0.33	0.47	0	1
zloc_eu	516	0.66	0.48	0	1
zloc_other	516	0.02	0.12	0	1
<b>Methodological choices</b>					
lca_att	516	0.98	0.14	0	1
lca_cons	516	0.02	0.14	0	1
wtt	516	0.34	0.47	0	1
wtw	516	0.66	0.47	0	1
infrastruct	516	0.43	0.50	0	1
copval_alloc_ener	516	0.28	0.45	0	1
copval_alloc_mass	516	0.05	0.21	0	1
copval_alloc_markval	516	0.13	0.34	0	1
copval_alloc_exerg	516	0.01	0.08	0	1
copval_alloc	516	0.47	0.50	0	1
copval_systexp	516	0.50	0.50	0	1
copval_hyb	516	0.00	0.04	0	1
gas_num	317	0.21	0.41	0	1
ass_ipcc	496	0.35	0.48	0	1
luc	516	0.58	0.49	0	1
luc_dir	516	0.58	0.49	0	1
luc_indir	516	0.03	0.16	0	1
uncer_ref	516	0.60	0.49	0	1
uncer_MC	516	0.11	0.32	0	1
uncer_SA	516	0.34	0.48	0	1
uncer_MCSA	516	0.46	0.50	0	1
impcat_nev	516	0.31	0.46	0	1
impcat_nrc	516	0.41	0.49	0	1
impcat_nrcnev	516	0.47	0.50	0	1
impcat_gwp	516	1.00	0.00	1	1
impcat_other	516	1.16	2.40	0	9
impcat_all	516	2.87	2.55	1	12
<b>Typology of the study</b>					
lit_pr	516	0.61	0.49	0	1
lit_or	516	0.10	0.30	0	1
lit_dir	516	0.02	0.14	0	1
lit_or_dir	516	0.12	0.33	0	1
lit_wp	516	0.26	0.44	0	1
year	516	2009.30	1.38	2002	2011
year_07	516	0.97	0.18	0	1
year_09	516	0.89	0.32	0	1
year_10	516	0.51	0.50	0	1
zlab_us	516	0.31	0.46	0	1
zlab_eu	516	0.69	0.46	0	1
zlab_other	516	0.00	0.00	0	0

**Table B.7 – Descriptive statistics for the effect size and variables of the original G2-Ethanol sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	361	29.45	48.39	-113.60	518.40
es_min	361	-2.17	51.44	-307.59	195.45
es_max	361	61.12	85.31	-49.54	841.35
es_var	361	941.37	4232.25	0.00	27149.13
es_et	361	16.19	26.10	0.03	164.77
<b>Technical data</b>					
gen_2	361	1.00	0.00	1	1
gen_3	361	0.00	0.00	0	0
etha	361	1.00	0.00	1	1
btI	361	0.00	0.00	0	0
fame	361	0.00	0.00	0	0
hvo	361	0.00	0.00	0	0
mat_algae	361	0.00	0.00	0	0
mat_agrires	361	0.29	0.45	0	1
mat_for	361	0.02	0.13	0	1
mat_enercult	361	0.29	0.45	0	1
mat_farmwood	361	0.10	0.30	0	1
mat_other	361	0.31	0.46	0	1
mat_cult	361	0.39	0.49	0	1
coprod	361	0.89	0.31	0	1
cop_gly	361	0.00	0.00	0	0
cop_elec	361	0.58	0.49	0	1
cop_heat	361	0.02	0.16	0	1
cop_algmeal	361	0.00	0.00	0	0
cop_biog	361	0.30	0.46	0	1
cop_other	361	0.15	0.36	0	1
cop_otherelec	361	0.46	0.50	0	1
coprod_num	361	1.06	0.49	0	3
mass_yield_exist	361	0.66	0.48	0	1
g2_mass_yield	237	0.25	0.05	0.11	0.50
eth_expl	361	0.08	0.28	0	1
eth_ac	361	0.83	0.37	0	1
eth_amm	361	0.08	0.28	0	1
eth_other	361	0.00	0.00	0	0
zloc_us	361	0.37	0.48	0	1
zloc_eu	361	0.61	0.49	0	1
zloc_other	361	0.02	0.15	0	1
<b>Methodological choices</b>					
lca_att	361	0.98	0.14	0	1
lca_cons	361	0.02	0.14	0	1
wtt	361	0.37	0.48	0	1
wtw	361	0.63	0.48	0	1
infrastructure	361	0.45	0.50	0	1
copval_alloc_ener	361	0.34	0.48	0	1
copval_alloc_mass	361	0.04	0.21	0	1
copval_alloc_markval	361	0.07	0.25	0	1
copval_alloc_exerg	361	0.01	0.09	0	1
copval_alloc	361	0.47	0.50	0	1
copval_systexp	361	0.52	0.50	0	1
copval_hyb	361	0.00	0.05	0	1
gas_num	205	0.12	0.33	0	1
ass_ipcc	361	0.35	0.48	0	1
luc	361	0.66	0.48	0	1
luc_dir	361	0.66	0.48	0	1
luc_indir	361	0.03	0.16	0	1
uncer_ref	361	0.54	0.50	0	1
uncer_MC	361	0.13	0.33	0	1
uncer_SA	361	0.38	0.49	0	1
uncer_MCSA	361	0.51	0.50	0	1
impcat_nev	361	0.20	0.40	0	1
impcat_nrc	361	0.34	0.47	0	1
impcat_nrcnev	361	0.37	0.48	0	1
impcat_gwp	361	1.00	0.00	1	1
impcat_other	361	0.89	2.04	0	9
impcat_all	361	2.43	2.11	1	11
<b>Typology of the study</b>					
lit_pr	361	0.62	0.49	0	1
lit_or	361	0.03	0.18	0	1
lit_dir	361	0.01	0.12	0	1
lit_ordir	361	0.05	0.21	0	1
lit_wp	361	0.33	0.47	0	1
year	361	2009.42	1.25	2002	2011
year_07	361	0.96	0.19	0	1
year_09	361	0.94	0.23	0	1
year_10	361	0.48	0.50	0	1
zlab_us	361	0.34	0.48	0	1
zlab_eu	361	0.66	0.48	0	1
zlab_other	361	0.00	0.00	0	0

**Table B.8 – Descriptive statistics for the effect size and variables of the original G2-BtL sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	155	19.04	35.78	-142.18	189.00
es_min	155	-14.77	40.94	-204.37	65.53
es_max	155	52.91	53.94	-127.01	334.35
es_var	155	560.76	975.95	0.21	5499.18
es_et	155	17.35	16.17	0.46	74.16
<b>Technical data</b>					
gen_2	155	1.00	0.00	1	1
gen_3	155	0.00	0.00	0	0
etha	155	0.00	0.00	0	0
btl	155	1.00	0.00	1	1
fame	155	0.00	0.00	0	0
hvo	155	0.00	0.00	0	0
mat_algae	155	0.00	0.00	0	0
mat_agrires	155	0.16	0.37	0	1
mat_for	155	0.15	0.36	0	1
mat_enercult	155	0.41	0.49	0	1
mat_farmwood	155	0.13	0.34	0	1
mat_other	155	0.15	0.36	0	1
mat_cult	155	0.54	0.50	0	1
coprod	155	0.90	0.31	0	1
cop_gly	155	0.00	0.00	0	0
cop_elec	155	0.93	0.26	0	1
cop_heat	155	0.20	0.40	0	1
cop_algmeal	155	0.00	0.00	0	0
cop_biog	155	0.00	0.00	0	0
cop_other	155	0.13	0.34	0	1
cop_otherelec	155	0.33	0.47	0	1
coprod_num	155	1.26	0.56	0	2
mass_yield_exist	155	0.91	0.29	0	1
g2_mass_yield	141	0.16	0.06	0.07	0.42
btl_pre_torr	153	0.07	0.25	0	1
btl_pre_pyro	153	0.05	0.21	0	1
btl_pre_none	153	0.89	0.32	0	1
btl_pro_utoth	153	0.86	0.35	0	1
btl_pro_alng	153	0.08	0.27	0	1
btl_pro_alelec	153	0.03	0.18	0	1
btl_pro_alrenew	153	0.03	0.16	0	1
btl_gasrecycl	153	0.01	0.11	0	1
btl_ccs	153	0.02	0.14	0	1
zloc_us	155	0.23	0.42	0	1
zloc_eu	155	0.77	0.42	0	1
zloc_other	155	0.00	0.00	0	0
<b>Methodological choices</b>					
lca_att	155	0.97	0.16	0	1
lca_cons	155	0.03	0.16	0	1
wtl	155	0.27	0.45	0	1
wtw	155	0.73	0.45	0	1
infrastruct	155	0.38	0.49	0	1
copval_alloc_ener	155	0.14	0.35	0	1
copval_alloc_mass	155	0.06	0.23	0	1
copval_alloc_markval	155	0.28	0.45	0	1
copval_alloc_exerg	155	0.00	0.00	0	0
copval_alloc	155	0.48	0.50	0	1
copval_systexp	155	0.46	0.50	0	1
copval_hyb	155	0.00	0.00	0	0
gas_num	112	0.38	0.49	0	1
ass_ipcc	135	0.34	0.48	0	1
luc	155	0.39	0.49	0	1
luc_dir	155	0.39	0.49	0	1
luc_indir	155	0.03	0.16	0	1
uncer_ref	155	0.72	0.45	0	1
uncer_MC	155	0.08	0.27	0	1
uncer_SA	155	0.26	0.44	0	1
uncer_MCSA	155	0.34	0.47	0	1
impcat_nev	155	0.56	0.50	0	1
impcat_nrc	155	0.57	0.50	0	1
impcat_nrcnev	155	0.71	0.46	0	1
impcat_gwp	155	1.00	0.00	1	1
impcat_other	155	1.79	2.99	0	9
impcat_all	155	3.92	3.11	1	12
<b>Typology of the study</b>					
lit_pr	155	0.59	0.49	0	1
lit_or	155	0.26	0.44	0	1
lit_dir	155	0.04	0.19	0	1
lit_ordir	155	0.30	0.46	0	1
lit_wp	155	0.11	0.31	0	1
year	155	2009.01	1.62	2002	2011
year_07	155	0.97	0.18	0	1
year_09	155	0.75	0.44	0	1
year_10	155	0.56	0.50	0	1
zlab_us	155	0.23	0.42	0	1
zlab_eu	155	0.77	0.42	0	1
zlab_other	155	0.00	0.00	0	0

## Appendix C. Complements on results

Table C.1 – Descriptive statistics for the effect size and variables of the econometric Whole sample

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	533	28.64	44.32	-85.00	332.20
es_min	533	-9.66	52.28	-352.81	101.59
es_max	533	66.99	105.24	-19.10	828.67
es_var	533	1666.43	8552.01	0.00	64159.87
es_et	533	19.61	35.84	0.03	253.30
<b>Technical data</b>					
gen_2	533	0.87	0.34	0	1
gen_3	533	0.13	0.34	0	1
etha	533	0.60	0.49	0	1
btl	533	0.27	0.44	0	1
fame	533	0.10	0.30	0	1
hvo	533	0.03	0.17	0	1
mat_algae	533	0.13	0.34	0	1
mat_agrires	533	0.20	0.40	0	1
mat_for	533	0.05	0.22	0	1
mat_enercult	533	0.30	0.46	0	1
mat_farmwood	533	0.10	0.30	0	1
mat_other	533	0.23	0.42	0	1
mat_cult	533	0.39	0.49	0	1
coprod	533	0.91	0.29	0	1
cop_gly	533	0.09	0.28	0	1
cop_elec	533	0.65	0.48	0	1
cop_heat	533	0.11	0.32	0	1
cop_algmeal	533	0.10	0.30	0	1
cop_biog	533	0.18	0.38	0	1
cop_other	533	0.24	0.53	0	2
cop_otherelec	533	0.46	0.50	0	1
coprod_num	533	1.35	0.99	0	5
mass_yield_exist	533	0.65	0.48	0	1
g2_mass_yield	341	0.22	0.07	0.07	0.50
eth_expl	321	0.08	0.27	0	1
eth_ac	321	0.83	0.37	0	1
eth_amm	321	0.09	0.28	0	1
eth_other	321	0.00	0.00	0	0
btl_pre_torr	141	0.06	0.23	0	1
btl_pre_pyro	141	0.04	0.20	0	1
btl_pre_none	141	0.90	0.30	0	1
btl_pro_autoth	141	0.88	0.33	0	1
btl_pro_alng	141	0.09	0.28	0	1
btl_pro_alelec	141	0.01	0.08	0	1
btl_pro_alrenew	141	0.03	0.17	0	1
btl_gasrecycl	141	0.01	0.08	0	1
btl_ccs	141	0.00	0.00	0	0
g3_productivity	68	31.05	19.58	6.85	150.00
g3_oil	69	0.41	0.11	0.15	0.70
g3_Oppond	69	0.68	0.47	0	1
zloc_us	533	0.32	0.47	0	1
zloc_eu	533	0.65	0.48	0	1
zloc_other	533	0.03	0.16	0	1
<b>Methodological choices</b>					
lca_att	533	0.97	0.18	0	1
lca_cons	533	0.03	0.18	0	1
wtt	533	0.36	0.48	0	1
wtw	533	0.64	0.48	0	1
infrastruct	533	0.42	0.49	0	1
copval_alloc_ener	533	0.27	0.44	0	1
copval_alloc_mass	533	0.05	0.21	0	1
copval_alloc_markval	533	0.10	0.30	0	1
copval_alloc_exerg	533	0.01	0.07	0	1
copval_alloc	533	0.42	0.49	0	1
copval_systexp	533	0.55	0.50	0	1
copval_hyb	533	0.07	0.26	0	1
wtw*g3_carbneut	69	0.00	0.00	0	0
gas_num	349	0.26	0.44	0	1
ass_ipcc	514	0.38	0.49	0	1
luc	533	0.52	0.50	0	1
luc_dir	533	0.52	0.50	0	1
luc_indir	533	0.04	0.19	0	1
uncer_ref	533	0.58	0.49	0	1
uncer_MC	533	0.10	0.29	0	1
uncer_SA	533	0.37	0.48	0	1
uncer_MCSA	533	0.47	0.50	0	1
impcat_nev	533	0.32	0.47	0	1
impcat_nrc	533	0.43	0.50	0	1
impcat_nrcnev	533	0.51	0.50	0	1
impcat_gwp	533	1.00	0.00	1	1
impcat_other	533	0.99	2.17	0	9
impcat_all	533	2.74	2.35	1	12
<b>Typology of the study</b>					
lit_pr	533	0.62	0.49	0	1
lit_or	533	0.09	0.29	0	1
lit_dir	533	0.03	0.18	0	1
lit_ordir	533	0.12	0.33	0	1
lit_wp	533	0.26	0.44	0	1
year	533	2009.39	1.32	2002	2011
year_07	533	0.97	0.18	0	1
year_09	533	0.90	0.30	0	1
year_10	533	0.58	0.49	0	1
zlab_us	533	0.32	0.47	0	1
zlab_eu	533	0.67	0.47	0	1
zlab_other	533	0.01	0.11	0	1

**Table C.2 – Descriptive statistics for the effect size and variables of the econometric G3 sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	69	58.84	104.99	-85.00	332.20
es_min	69	-58.50	91.73	-352.81	101.59
es_max	69	176.17	243.48	3.92	828.67
es_var	69	9439.33	21386.86	265.09	64159.87
es_et	69	59.86	77.08	16.28	253.30
<b>Technical data</b>					
gen_2	69	0.00	0.00	0	0
gen_3	69	1.00	0.00	1	1
etha	69	0.00	0.00	0	0
btl	69	0.00	0.00	0	0
fame	69	0.80	0.41	0	1
hvo	69	0.22	0.42	0	1
mat_algae	69	1.00	0.00	1	1
mat_agrires	69	0.00	0.00	0	0
mat_for	69	0.00	0.00	0	0
mat_enercult	69	0.00	0.00	0	0
mat_farmwood	69	0.00	0.00	0	0
mat_other	69	0.00	0.00	0	0
mat_cult	69	0.00	0.00	0	0
coprod	69	1.00	0.00	1	1
cop_gly	69	0.68	0.47	0	1
cop_elec	69	0.30	0.46	0	1
cop_heat	69	0.39	0.49	0	1
cop_algmeal	69	0.77	0.43	0	1
cop_biog	69	0.00	0.00	0	0
cop_other	69	0.94	0.92	0	2
cop_otherelec	69	0.86	0.35	0	1
coprod_num	69	2.93	1.72	1	5
mass_yield_exist	69	0.06	0.24	0	1
g3_productivity	68	31.05	19.58	6.85	150.00
g3_oil	69	0.41	0.11	0.15	0.70
g3_Oppond	69	0.68	0.47	0	1
zloc_us	69	0.41	0.49	0	1
zloc_eu	69	0.51	0.50	0	1
zloc_other	69	0.09	0.28	0	1
<b>Methodological choices</b>					
lca_att	69	0.91	0.28	0	1
lca_cons	69	0.09	0.28	0	1
wtw	69	0.61	0.49	0	1
wtw	69	0.39	0.49	0	1
infrastruct	69	0.51	0.50	0	1
copval_alloc_ener	69	0.09	0.28	0	1
copval_alloc_mass	69	0.00	0.00	0	0
copval_alloc_markval	69	0.03	0.17	0	1
copval_alloc_exerg	69	0.00	0.00	0	0
copval_alloc	69	0.12	0.32	0	1
copval_systexp	69	0.88	0.32	0	1
copval_hyb	69	0.55	0.50	0	1
wtw*g3_carbneut	69	0.00	0.00	0	0
gas_num	69	0.45	0.50	0	1
ass_ipcc	69	0.48	0.50	0	1
luc	69	0.09	0.28	0	1
luc_dir	69	0.09	0.28	0	1
luc_indir	69	0.09	0.28	0	1
uncer_ref	69	0.43	0.50	0	1
uncer_MC	69	0.00	0.00	0	0
uncer_SA	69	0.57	0.50	0	1
uncer_MCSA	69	0.57	0.50	0	1
impcat_nev	69	0.28	0.45	0	1
impcat_nrc	69	0.39	0.49	0	1
impcat_nrcnev	69	0.67	0.47	0	1
impcat_gwp	69	1.00	0.00	1	1
impcat_other	69	0.68	1.18	0	5
impcat_all	69	2.35	1.44	1	7
<b>Typology of the study</b>					
lit_pr	69	0.70	0.46	0	1
lit_or	69	0.00	0.00	0	0
lit_dir	69	0.09	0.28	0	1
lit_ordir	69	0.09	0.28	0	1
lit_wp	69	0.22	0.42	0	1
year	69	2009.87	0.34	2009	2010
year_07	69	1.00	0.00	1	1
year_09	69	1.00	0.00	1	1
year_10	69	0.87	0.34	0	1
zlab_us	69	0.41	0.49	0	1
zlab_eu	69	0.51	0.50	0	1
zlab_other	69	0.09	0.28	0	1

**Table C.3 – Descriptive statistics for the effect size and variables of the econometric G2 sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	464	24.15	21.95	-24.00	85.80
es_min	464	-2.40	38.69	-307.59	49.94
es_max	464	50.75	44.47	-19.10	380.55
es_var	464	510.55	2552.67	0.00	27149.13
es_et	464	13.62	18.05	0.03	164.77
<b>Technical data</b>					
gen_2	464	1.00	0.00	1	1
gen_3	464	0.00	0.00	0	0
etha	464	0.69	0.46	0	1
btI	464	0.31	0.46	0	1
fame	464	0.00	0.00	0	0
hvo	464	0.00	0.00	0	0
mat_algae	464	0.00	0.00	0	0
mat_agríres	464	0.23	0.42	0	1
mat_for	464	0.06	0.23	0	1
mat_enercult	464	0.34	0.47	0	1
mat_farmwood	464	0.11	0.32	0	1
mat_other	464	0.26	0.44	0	1
mat_cult	464	0.45	0.50	0	1
coprod	464	0.89	0.31	0	1
cop_gly	464	0.00	0.00	0	0
cop_elec	464	0.70	0.46	0	1
cop_heat	464	0.07	0.26	0	1
cop_algmeal	464	0.00	0.00	0	0
cop_biog	464	0.21	0.41	0	1
cop_other	464	0.14	0.35	0	1
cop_otherelec	464	0.40	0.49	0	1
coprod_num	464	1.12	0.51	0	3
mass_yield_exist	464	0.73	0.44	0	1
g2_mass_yield	341	0.22	0.07	0.07	0.50
eth_expl	321	0.08	0.27	0	1
eth_ac	321	0.83	0.37	0	1
eth_amm	321	0.09	0.28	0	1
eth_other	321	0.00	0.00	0	0
btI_pre_torr	141	0.06	0.23	0	1
btI_pre_pyro	141	0.04	0.20	0	1
btI_pre_none	141	0.90	0.30	0	1
btI_pro_autoth	141	0.88	0.33	0	1
btI_pro_alng	141	0.09	0.28	0	1
btI_pro_alelec	141	0.01	0.08	0	1
btI_pro_alrenew	141	0.03	0.17	0	1
btI_gasrecycl	141	0.01	0.08	0	1
btI_ccs	141	0.00	0.00	0	0
zloc_us	464	0.31	0.46	0	1
zloc_eu	464	0.67	0.47	0	1
zloc_other	464	0.02	0.13	0	1
<b>Methodological choices</b>					
lca_att	464	0.98	0.15	0	1
lca_cons	464	0.02	0.15	0	1
wtt	464	0.32	0.47	0	1
wtw	464	0.68	0.47	0	1
infrastruct	464	0.40	0.49	0	1
copval_alloc_ener	464	0.29	0.46	0	1
copval_alloc_mass	464	0.05	0.23	0	1
copval_alloc_markval	464	0.11	0.32	0	1
copval_alloc_exerg	464	0.01	0.08	0	1
copval_alloc	464	0.47	0.50	0	1
copval_systexp	464	0.50	0.50	0	1
copval_hyb	464	0.00	0.05	0	1
gas_num	280	0.21	0.41	0	1
ass_ipcc	445	0.36	0.48	0	1
luc	464	0.58	0.49	0	1
luc_dir	464	0.58	0.49	0	1
luc_indir	464	0.03	0.17	0	1
uncer_ref	464	0.60	0.49	0	1
uncer_MC	464	0.11	0.31	0	1
uncer_SA	464	0.34	0.47	0	1
uncer_MCSA	464	0.45	0.50	0	1
impcat_nev	464	0.33	0.47	0	1
impcat_nrc	464	0.43	0.50	0	1
impcat_nrcnev	464	0.49	0.50	0	1
impcat_gwp	464	1.00	0.00	1	1
impcat_other	464	1.04	2.28	0	9
impcat_all	464	2.80	2.45	1	12
<b>Typology of the study</b>					
lit_pr	464	0.61	0.49	0	1
lit_or	464	0.10	0.30	0	1
lit_dir	464	0.02	0.15	0	1
lit_ordir	464	0.13	0.33	0	1
lit_wp	464	0.26	0.44	0	1
year	464	2009.32	1.40	2002	2011
year_07	464	0.96	0.19	0	1
year_09	464	0.89	0.32	0	1
year_10	464	0.53	0.50	0	1
zlab_us	464	0.30	0.46	0	1
zlab_eu	464	0.70	0.46	0	1
zlab_other	464	0.00	0.00	0	0

**Table C.4 – Descriptive statistics for the effect size and variables of the econometric G2-Ethanol sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	321	26.61	22.26	-23.65	85.80
es_min	321	1.81	41.79	-307.59	49.94
es_max	321	51.45	45.93	-14.10	380.55
es_var	321	532.67	3020.32	0.00	27149.13
es_et	321	12.72	19.29	0.03	164.77
<b>Technical data</b>					
gen_2	321	1.00	0.00	1	1
gen_3	321	0.00	0.00	0	0
etha	321	1.00	0.00	1	1
btl	321	0.00	0.00	0	0
fame	321	0.00	0.00	0	0
hvo	321	0.00	0.00	0	0
mat_algae	321	0.00	0.00	0	0
mat_agrires	321	0.26	0.44	0	1
mat_for	321	0.02	0.14	0	1
mat_enercult	321	0.31	0.46	0	1
mat_farmwood	321	0.10	0.30	0	1
mat_other	321	0.31	0.46	0	1
mat_cult	321	0.41	0.49	0	1
coprod	321	0.89	0.31	0	1
cop_gly	321	0.00	0.00	0	0
cop_elec	321	0.60	0.49	0	1
cop_heat	321	0.02	0.15	0	1
cop_algmeal	321	0.00	0.00	0	0
cop_biog	321	0.30	0.46	0	1
cop_other	321	0.14	0.35	0	1
cop_otherelec	321	0.44	0.50	0	1
coprod_num	321	1.07	0.48	0	3
mass_yield_exist	321	0.65	0.48	0	1
g2_mass_yield	209	0.25	0.05	0.14	0.50
eth_expl	321	0.08	0.27	0	1
eth_ac	321	0.83	0.37	0	1
eth_amm	321	0.09	0.28	0	1
eth_other	321	0.00	0.00	0	0
zloc_us	321	0.35	0.48	0	1
zloc_eu	321	0.63	0.48	0	1
zloc_other	321	0.02	0.16	0	1
<b>Methodological choices</b>					
lca_att	321	0.98	0.15	0	1
lca_cons	321	0.02	0.15	0	1
wt	321	0.36	0.48	0	1
wtw	321	0.64	0.48	0	1
infrastruct	321	0.42	0.49	0	1
copval_alloc_ener	321	0.36	0.48	0	1
copval_alloc_mass	321	0.05	0.22	0	1
copval_alloc_markval	321	0.05	0.22	0	1
copval_alloc_exerg	321	0.01	0.10	0	1
copval_alloc	321	0.47	0.50	0	1
copval_systexp	321	0.50	0.50	0	1
copval_hyb	321	0.00	0.06	0	1
gas_num	177	0.12	0.33	0	1
ass_ipcc	321	0.37	0.48	0	1
luc	321	0.66	0.47	0	1
luc_dir	321	0.66	0.47	0	1
luc_indir	321	0.03	0.17	0	1
uncer_ref	321	0.53	0.50	0	1
uncer_MC	321	0.12	0.33	0	1
uncer_SA	321	0.39	0.49	0	1
uncer_MCSA	321	0.52	0.50	0	1
impcat_nev	321	0.22	0.42	0	1
impcat_nrc	321	0.36	0.48	0	1
impcat_nrcnev	321	0.39	0.49	0	1
impcat_gwp	321	1.00	0.00	1	1
impcat_other	321	0.73	1.81	0	9
impcat_all	321	2.32	1.92	1	11
<b>Typology of the study</b>					
lit_pr	321	0.61	0.49	0	1
lit_or	321	0.04	0.19	0	1
lit_dir	321	0.02	0.12	0	1
lit_ordir	321	0.05	0.22	0	1
lit_wp	321	0.33	0.47	0	1
year	321	2009.45	1.26	2002	2011
year_07	321	0.96	0.19	0	1
year_09	321	0.94	0.24	0	1
year_10	321	0.51	0.50	0	1
zlab_us	321	0.34	0.47	0	1
zlab_eu	321	0.66	0.47	0	1
zlab_other	321	0.00	0.00	0	0

**Table C.5 – Descriptive statistics for the effect size and variables of the econometric G2-BtL sample**

	# of values	mean	std dev	min	max
<b>Descriptive statistics for the Effect Size</b>					
es	143	18.65	20.25	-24.00	85.68
es_min	143	-11.84	28.54	-120.95	43.58
es_max	143	49.19	41.12	-19.10	169.75
es_var	143	460.89	827.83	0.21	5499.18
es_et	143	15.66	14.74	0.46	74.16
<b>Technical data</b>					
gen_2	143	1.00	0.00	1	1
gen_3	143	0.00	0.00	0	0
etha	143	0.00	0.00	0	0
btl	143	1.00	0.00	1	1
fame	143	0.00	0.00	0	0
hvo	143	0.00	0.00	0	0
mat_algae	143	0.00	0.00	0	0
mat_agrires	143	0.15	0.36	0	1
mat_for	143	0.15	0.36	0	1
mat_enercult	143	0.41	0.49	0	1
mat_farmwood	143	0.13	0.34	0	1
mat_other	143	0.15	0.36	0	1
mat_cult	143	0.55	0.50	0	1
coprod	143	0.89	0.32	0	1
cop_gly	143	0.00	0.00	0	0
cop_elec	143	0.92	0.27	0	1
cop_heat	143	0.18	0.39	0	1
cop_algmeal	143	0.00	0.00	0	0
cop_biog	143	0.00	0.00	0	0
cop_other	143	0.13	0.34	0	1
cop_otherelec	143	0.31	0.47	0	1
coprod_num	143	1.24	0.56	0	2
mass_yield_exist	143	0.92	0.27	0	1
g2_mass_yield	132	0.16	0.06	0.07	0.42
btl_pre_torr	141	0.06	0.23	0	1
btl_pre_pyro	141	0.04	0.20	0	1
btl_pre_none	141	0.90	0.30	0	1
btl_pro_autoth	141	0.88	0.33	0	1
btl_pro_alng	141	0.09	0.28	0	1
btl_pro_alelec	141	0.01	0.08	0	1
btl_pro_alrenew	141	0.03	0.17	0	1
btl_gasrecycl	141	0.01	0.08	0	1
btl_ccs	141	0.00	0.00	0	0
zloc_us	143	0.22	0.42	0	1
zloc_eu	143	0.78	0.42	0	1
zloc_other	143	0.00	0.00	0	0
<b>Methodological choices</b>					
lca_att	143	0.97	0.17	0	1
lca_cons	143	0.03	0.17	0	1
wt	143	0.24	0.43	0	1
wtw	143	0.76	0.43	0	1
infrastruct	143	0.36	0.48	0	1
copval_alloc_ener	143	0.14	0.35	0	1
copval_alloc_mass	143	0.06	0.24	0	1
copval_alloc_markval	143	0.25	0.44	0	1
copval_alloc_exerg	143	0.00	0.00	0	0
copval_alloc	143	0.45	0.50	0	1
copval_systexp	143	0.48	0.50	0	1
copval_hyb	143	0.00	0.00	0	0
gas_num	103	0.36	0.48	0	1
ass_ipcc	124	0.35	0.48	0	1
luc	143	0.41	0.49	0	1
luc_dir	143	0.41	0.49	0	1
luc_indir	143	0.03	0.17	0	1
uncer_ref	143	0.76	0.43	0	1
uncer_MC	143	0.08	0.27	0	1
uncer_SA	143	0.22	0.42	0	1
uncer_MCSA	143	0.30	0.46	0	1
impcat_nev	143	0.57	0.50	0	1
impcat_nrc	143	0.59	0.49	0	1
impcat_nrcnev	143	0.71	0.45	0	1
impcat_gwp	143	1.00	0.00	1	1
impcat_other	143	1.73	2.99	0	9
impcat_all	143	3.90	3.10	1	12
<b>Typology of the study</b>					
lit_pr	143	0.61	0.49	0	1
lit_or	143	0.25	0.44	0	1
lit_dir	143	0.04	0.20	0	1
lit_ordir	143	0.29	0.46	0	1
lit_wp	143	0.10	0.30	0	1
year	143	2009.05	1.63	2002	2011
year_07	143	0.97	0.18	0	1
year_09	143	0.76	0.43	0	1
year_10	143	0.58	0.50	0	1
zlab_us	143	0.22	0.42	0	1
zlab_eu	143	0.78	0.42	0	1
zlab_other	143	0.00	0.00	0	0

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