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Concentration Indices of Income related Self-Reported Health: A Meta-Regression Analysis

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Abstract

Proliferating evidence reporting on standardised cross-country concentration indexes of income related self-reported health is increasingly being used for policy evaluation. Nonetheless, limited efforts have been put forward to examine the extent to which such evidence is subject to any specific methodological and publication biases, given that studies relying upon survey data form different samples and both heterogeneous health system institutions and empirical strategies. We conduct the first study drawing upon appropriate statistical methods to examine the presence of publication bias in the health economics literature measuring health inequalities of self-reported health. We test for other biases including the effect of precision estimates based on meta-regression analysis (MRA). We account for a set of biases in estimates of income-related health inequalities that rely on concentration index-related methods and self-reported health measures. Our findings suggest evidence of publication bias that primarily depends on the cardinalisation of self-reported health and some evidence of study-specific precision.

Keywords: health inequalities, concentration index, self-reported health, publication bias
1. Introduction

Health inequalities are generally regarded as a key outcome measure in order to evaluate health system performance worldwide (WHO, 2000). WHO estimates indicate the existence of gaps in health outcomes not only across countries but especially within countries, which do not appear to fade away on time (Costa-Font et al, 2011). The latter are rooted in pervasive differences in social status, income, ethnicity, and gender, among other determinants that appear to impact on health production. Some studies document that the health of a 20-year-old low-income male, on average, reports to be in similar health as a 60-year-old high-income male in the United States (Case and Deaton, 2005). Efforts to reduce health inequalities together as improving health outcomes make the most part of health and healthcare policies worldwide.

In response to such as policy need, recent health policy research has focused on developing sound methodologies to undertake such measurements, primarily drawing from index measures that meet some ideal properties, and more specifically, concentration indices. Concentration indices are a measure of health inequality that can be transformed to measure inequality in the distribution of health (Wagstaff, 2002). The advantage of such a measure is that summarizes a large range of distributional information, which is relatively simple to compute, transform, easy, graph and interpret. As we explain below, the index is computed by ranking individuals based on socio-economic status, and computing the cumulative proportions of the population (beginning with the most disadvantaged and ending with the least disadvantaged) against the cumulative proportion of health.

Nonetheless, it is a mean dependent index, and hence changes in the population mean will influence the inequality estimates. Second, as an index it assumes that the variable to which the index is applied adopts a continuous dimension. In measuring health, analysts generally follow some form of cardinalisation strategy, or rely on ordinal or interval regression to specifically transform categorical health data into a continuous fashion. However, limited analysis has been undertaken on the empirical performance of concentration index measures in health. Most of the literature focuses on the value judgments adopted by the methodology and the potential biases that concentration indices exert in inequality measurement.

The wealth of evidence on heterogeneity in existing estimates suggests that there are reasons to believe that publication bias exists. Often studies rely on different datasets of similar European
countries, use different inferences and often carry out adjustments to adequate the measure of self-reported health employed to ideal requirements (Van Doorslaer et al, 1997, 2004). However, limited meta-analysis, or meta-regression studies have been undertaken to account for the numerous study biases that are generally present in the empirical literature, and that we ascertain are not absent from health inequality studies. The health economics literature is a prone area for biased estimates (Costa-Font et al, 2013 for a review). One of the areas where biased estimates can emerge is in the measuring of health inequalities due to the large difficulties in measuring health, accounting for study and institutional constraints, as well as study year and data alongside other potential explanations for publication bias such as precision. There are plausible reasons to assume that this can be the case in clinical trials and experimental studies, this assumption do not necessarily hold for studies based on survey data. Analyses of survey data that measure inequality using self-reported health can rarely rule out unobserved heterogeneity as a source of bias. Hence, we inquire about how important are all those potential biases in explaining the heterogeneity in health inequality estimates.

This paper attempts to examine the extent to which income related inequalities in health are affected by precision and publication biases by drawing on MRA. Our contribution to the literature is to identify the existence of biases in studies measuring socioeconomic inequalities in health. More specifically, we focus in the predominant measure of health inequality, namely concentration index estimates of self reported health, which is the main measure of health outcome employed ain the health economic literature. We examine precision effects, the sort of publication outlets they get published on, alongside other study characteristics that could potentially shape the empirical estimates in some direction. In doing so, it is then possible to use the meta-regression analysis (MRA) - a set of techniques developed to integrate and correct estimated regression coefficients. Thus, each country will have a "true" CI in any given year and pooling estimates that employ the same methodology them together and identifying what factors explain their heterogeneity, we claim allows filtering the sort of biases, and hence coming up with an unbiased estimate that nets outs the specific country, system and study heterogeneity etc. A second objective lies in explaining the determinants of health inequality estimates taking advantage of MRA. Indeed, MRA produces estimates after correcting for precision effects (generally proxied by the standard error of the estimates). In addition, such regression can incorporate institutional determinants of the countries (e.g., whether countries are national health systems (NHS) and hence tax financed), which can explain inequality estimates. More specifically, we have tested for the existence of different biases that explain inequality estimates when study characteristics and methodologies or empirical strategies are controlled for. We rely on estimates that use concentration index to make sure we
examine and homogenous sample, which in turn used a measure of self-reported health that relies on a comparable temporal period.

Given the heterogeneity in inequality measurement methodologies in social science, and in the health status measures, we restrict our analysis to studies that employed homogeneous inequality indexes (generally representing the methods health economists rely on), and more specifically concentration indexes. Furthermore, given the distinct meaning of health status measures, we in addition restricted our sample to studies that employ measures of self–reported health. The empirical strategy followed is to first graphically examine funnel graphs, which plot estimates against a measure of precision. The latter is informative of the distribution of the sample of studies examined. Next, we undertake multivariate MRA to explain the typically large systematic variation among reported effects and estimate the size of potential biases. With sufficient data, we can sensibly estimate the effects that various methodological choices have upon the magnitude of the reported empirical results. It is important to acknowledge that a meta-analysis of observational studies has a number of potential limitations or pitfalls (Egger et al. 1997).

Generally speaking, meta-analysis are well-suited if the studies included produce average consistent, unbiased effect estimates. While there are plausible reasons to assume that this can be the case in clinical trials and experimental studies, this assumption does not necessarily hold for studies based on survey data. Analyses of survey data that measure inequality using self-reported health can rarely rule out unobserved heterogeneity as a source of bias. For this reason, we extend the MRA to control for a set of potential study and context specific characteristics we attempt to control for some of the unobserved heterogeneity.

To summarise, this paper aims at examining the following issues:

a) How country-specific determinants of health inequalities influence estimates of concentration indices of self-reported health
b) After controlling for health system specific effects, we attempt to to isolate the effect of precision effects in estimates of concentration index of self-reported health
c) In explaining the determinants of the heterogeneity in concentration index estimates, we aim at identifying some of the underpinning determinants where we know less about.

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1 A funnel graph is a scatter diagram of a reported empirical estimate (c) and its precision (1/SE).
The structure of the paper is as follows. In Section 2 we present the methods and data used for this analysis. In Section 3 we offer a discussion of the results and Section 4 is devoted to conclusions and implications.

2. Methods and Data

2.1 Methods and empirical strategy

Measuring inequalities in health

Inequality is in itself a measure of relative dispersion that can be identified visually by comparing extremes on a distribution. However, the measure encounters severe difficulties when it comes to finding ways to compare two country distributions over time and space. One way to summarise such information is by using inequality indices. Inequalities indices include ranking, Lorenz curves and Gini coefficients, concentration curves and concentration Indices.

The Lorenz Curve and Gini coefficient measure the absolute level of inequality in health (LeGrand, 1989; Wagstaff, Paci and van Doorslaer, 1991) and the expression is given by:

\[
G = \left(\frac{2}{\mu}\right) \text{Cov}(y, R_h)
\]

where \(R_h\) is the relative rank in the health distribution, with individuals ordered from the lowest to the highest level of health.

Similarly, concentration curves can be used to evaluate to what extent certain characteristics are unequally distributed according to health, not income, and to calculate the concentration indices.

There are three basic requirements of an inequity index: i) to reflect the socioeconomic dimension of inequalities in health, ii) to reflect the experiences of the population as a whole, and iii) to be sensitive to changes in the distribution of the population among socioeconomic groups. Many indices, such as the Gini coefficient, do not satisfy the first requirement. Others, such as ranking, do not take into account the other two: they only focus on the experience of the groups at the extreme of the distribution and they do not reflect the distribution of the population in several groups. The main advantages of Concentration Indices are that they meet the basic requirements and they are an easy way to compare inequalities among countries. In addition, they are useful for several reasons: to
check whether the relative magnitude in some country is important and to evaluate which health care systems contribute more to widening levels of inequality.

The Gini (G) coefficient and the Concentration index (CI) are directly related through the following expression (we have excluded sub-indexes for simplicity):

\[
CI = \left[ \frac{r(y,R_y)}{r(y,R_h)} \right]G \tag{2}
\]

Where \( \rho \) refers to the correlation coefficient, \( R_y \) refers to the ranking of socio-economic status and \( R_h \) the ranking of individuals based on their health status.

Policy makers may also be concerned about other sources of inequality that are captured in a measure of total health inequality. This can be analysed using health Lorenz curves and inequality can be measured using the Gini coefficient of health inequality (Le Grand, 1989; Wagstaff, Paci and van Doorslaer, 1991). The attraction of this approach is that there is a direct relationship between the concentration index and the Gini coefficient for health: the concentration index is proportional to the Gini coefficient, where the factor of proportionality is given by the ratio between the correlation coefficient for health and income rank and the correlation coefficient between health and health rank (Kakwani, 1980; van Doorslaer and Jones, 2003).

Inequality estimates based on concentration curves and concentration indices have been extensively used for measuring inequalities and inequities (Wagstaff and van Doorslaer, 2000). The health concentration curve (CC) and concentration index (CI) provide measures of relative income-related health inequality (Wagstaff, Van Doorslaer and Paci, 1989). Wagstaff, Paci and van Doorslaer (1991) have reviewed and compared the properties of the concentration curves and indices with alternative measures of health inequality. They argue the main advantages as the following: they capture the socioeconomic dimension of health inequalities; they use information from the whole income distribution rather than just the extremes; they provide the possibility to represent results visually through the concentration curve; and finally, they allow checks for dominance relationships.

The concentration index (CI) is derived from the concentration curve (CC). This is illustrated in Figure 1 for a measure of ill health. The sample of interest is ranked by socioeconomic status. If income is used as the relevant ranking variable, the horizontal axis begins with the poorest individual
and progresses through the income distribution up to the richest individual. This relative income rank is then plotted against the cumulative proportion of illness on the vertical axis. This assumes that a cardinal measure of illness is available that can be compared and aggregated across individuals. The 45-degree line shows the line of perfect equality, along which the population shares of illness are proportional to income, such that the poorest 20% of individuals experience 20% of illnesses in the population. “Pro-poor” inequality is illustrated by the concave curve in the figure, which corresponds to the concentration curve. In the example shown, the poorest 20% of income earners experience more than 20% of illnesses. The size of inequality can be summarised by the health concentration index, which is given by twice the area between the concentration curve and the 45-degree line.

Figure 1: Concentration curve for ill-health

Source: Authors’ elaboration

There are various ways of expressing the CI algebraically. The one that is mostly used in the literature for its convenience is:

\[
C = \frac{2}{\mu} \sum_{i=1}^{N} (y_i - \mu)(R_i - \frac{1}{2}) = \frac{2}{\mu} \text{cov}(y_i, R_i)
\]  

(3)
This shows that the value of the concentration index is equal to the covariance between individual health \( h_i \) and the individual’s relative rank \( R_i \), scaled by the mean of health in the population \( \mu \). Then to ensure the concentration index ranges between -1 and +1, the whole expression is multiplied by 2. Equation (1) indicates that the CI is a measure of the degree of association between an individual’s level of health and their relative position in the income distribution. It is important to highlight that a value of CI = 0 does not mean an absence of inequality, but an absence of the socioeconomic gradient in the distribution; this is, an absence of inequality associated with socioeconomic characteristics.

2.2 Meta-regression analysis

The standard MRA model used in the vast majority of economic applications is:

\[
e_j = \beta + \sum \alpha_k Z_{jk} + \varepsilon_j \quad (j=1, 2, \ldots L)
\]  

Where \( e_j \) is the empirical effect in question, and \( Z_{jk} \) are moderator variables used to explain the large study-to-study heterogeneity routinely found in economics research (Stanley and Jarrell, 1989). Moderator variables might include:

1. Dummy variables, which reflect whether potentially relevant independent variables have been omitted from (or included in) the primary study.
2. Specification variables that account for differences in functional form, types of regression, and data definitions or sources, etc.

The conventional model of publication selection in both economics and medical research is a simple MRA between a study’s estimated effect and its standard error.

\[
CI_i = \beta_1 + \beta_0S E_i + \varepsilon_i
\]  

\[(\text{Egger et al., 1997; Stanley, 2005; Stanley, 2008}).\]

An obvious statistical problem is that estimated effects in equation (5) will have different variances (i.e., heteroschedasticity). Weighted least squares (WLS) are the conventional correction for

---

2 As discussed in the next section, one of these moderator variables should be the estimate’s standard error if we are to identify and control for publication selection bias.
heteroschedasticity. The WLS version of (5) may be obtained by weighting the squared errors by the inverse of the estimates’ individual variances (i.e., \(1/SE_i^2\)), or by dividing equation (5) by \(SE_i\). In doing so, the resulting model is given by (6):

\[
t_i = \frac{CI_i}{SE_i} = \beta_0 + \beta_1 \left(\frac{1}{SE_i}\right) + \beta_2 X_i \left(\frac{1}{SE_i}\right) + v_i
\]

Note that the dependent variable becomes the study’s reported t-value, and the independent variable is precision, \(1/SE_i\). As \(SE_i\) approaches zero in equation (5), the expected effect will approach \(\beta_1\), regardless of publication selection bias. For this reason, medical researchers use the estimate of \(\beta_1\) in equation (5) or (6) as the corrected empirical effect. \(^4\) \(X_i\) refers to the set of other covariates that are study specific and are thought to influence the empirical estimates. Both the funnel graph and this MRA model of publication selection reveal the central importance of precision in evaluating research. Testing precision’s coefficient (\(H_0: \beta_1=0\)) serves as a powerful statistical test—precision-effect test (PET) — for a genuine empirical effect beyond publication selection (Stanley, 2008). PET’s validity has been confirmed in simulations and in several economic applications (Stanley, 2008; Doucouliagos and Stanley, 2009). The significance of the constant term is known as the FAT - funnel-asymmetry test.

Finally, as an extension, a Heckman-like correction called Precision effect estimate with standard error (PEESE) is provided, which refers to the precision effect estimate with standard error model, and can be used to obtain an estimate that is robust to publication selection bias so that (6) can be extended to:

\[
t_i = \frac{CI}{SE_i} = \beta_0 \cdot SE_i + \beta_1 \left(\frac{1}{SE_i}\right) + \beta_2 X_i \left(\frac{1}{SE_i}\right) + v_i
\]

2.3 Data

The data used in this study has been built by carefully reading and coding published studies identified in Medline, Econ lit and Sociofile. The search algorithm that is used to identify the relevant estimates of the literature was paper published in (a wide variety of journals) journals the

\[^3\] Rather than actually dividing all the observations of each variable by \(SE_i\), many meta-analysts choose to use a canned statistical routine for WLS using \(1/SE_i^2\) as the weights. Estimating equation (6) using OLS gives the same results as standard statistical routines for WLS on equation (5).

\[^4\] Unfortunately, this estimate is biased downward when there is a genuine nonzero effect (Stanley, 2008). To reduce this bias, Stanley and Doucouliagos (2007) offer an alternative MRA estimator. Also see Moreno et al. (2009).
time of the study ending in 2012 where the concentration index methodology was employed on self-reported health. Estimates where then examined manually and only those studies that reported a coefficient estimate and either the standard error (directly or indirectly) were kept. From the sample obtained we proceeded to selecting those that used a homogeneous measure of health that appears to be more prevalent, namely self-reported health status. When some of the information was not present in the study, we have inferred it from other paper estimates or asked authors to provide it so that a full database could be constructed. In some cases, we identified some errors in the original paper and we have corrected them in our estimate. From each study, we selected a set of relevant variables including: sample size, number of variables, method employed, institutional variables, precision and other relevant characteristics.

Table 1 reports the summary statistics of the main variables employed in the study. Specifically, our dependent variable is an estimate of the concentration index of self-reported health for a set of different countries (CI). Consistently with the literature, given that studies using categorical variables rely on transformation of self-reported health on measures of ill health estimated using the conventional scales, a negative concentration index is suggestive of ill health concentrated among the less affluent. However, we take the absolute value of significant estimates to ease the interpretation of the results. The average value of the concentration index is roughly 0.05, which exhibits a significant standard error (SE), suggesting the existence of significant heterogeneity in concentration index estimates, as exhibited in the Funnel plot. Furthermore, conventionally, MRA estimates include as covariates the standard error of each CI estimate (which proxies for the precision of each estimate) and exhibit a mean value of 0.015. Given that most studies supply European data, we have classified estimates based on some identifiable features of the health system, namely whether the data refers to a country where the health system is organised as a public national health service (NHS) (around 46% in our sample) or not. The latter is important so long as national health services tend to prioritise equity as a system goal under the mission of ‘equal access for equal need’. NHS is a dummy variable taking the value of 1 if an estimate refers to a set of countries in Northern Europe as well as a few in southern Europe including Britain and Ireland, whilst countries organised as social insurance schemes would take that value of zero. Then our study incorporates variables proxying the year of the study (Year), which arguably will influence both the magnitude and the precision of the inequality estimates given that inequality indexes often have been improved over time in part due to the increasing issue of panel data techniques which account for potential sources of unobserved heterogeneity. In addition, other controls that were deemed relevant were the number of observations (N) - the larger the number of estimates, the more reliable they are. Finally, given the complexity in
measuring health, we examine whether health status as a variable was cardinalised (Cardinal) which refers to 88% of the cases included in the analysis, or instead whether health was measured in an ordinal or categorical format. The dataset contain other variables that we did not use in the analysis as they turn out to be insignificant, such as the journal impact factor where the estimated was published, and the influence of the range of categories in using self-reported health. However, given that most estimates are based on manipulation of such variables controlling for such a variable shows no significant effects either and was finally dropped for the meta-regression analysis.

### Table 1. Summary Statistics

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
<th>Number of Observations</th>
<th>Mean (s.e)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CI</td>
<td>Concentration Index Estimate</td>
<td>301</td>
<td>0.048 (0.002)</td>
</tr>
<tr>
<td>SE</td>
<td>Standard error of the concentration index</td>
<td>298</td>
<td>0.015 (0.001)</td>
</tr>
<tr>
<td>NHS</td>
<td>Estimate from a National Health Service</td>
<td>301</td>
<td>0.465 (0.028)</td>
</tr>
<tr>
<td>Year</td>
<td>Year of the estimate - 1978</td>
<td>195</td>
<td>16.4 (0.424)</td>
</tr>
<tr>
<td>N</td>
<td>Number of observations</td>
<td>139</td>
<td>6399 (424.5)</td>
</tr>
<tr>
<td>Cardinal</td>
<td>Some form of cardinal transformation is performed</td>
<td>301</td>
<td>0.887 (0.018)</td>
</tr>
</tbody>
</table>

### 3. Results

After extracting estimates for all the studies identified in the sample, we were left with 301 observations, which constitute a sample of homogeneous observations very much in line with other meta-regression studies. Although new studies are being produced every year, the number of studies already meeting publication standards is sufficient to perform a meta-regression analysis, given that they draw upon methods developed about two decades ago.

Possibly the first and most natural way to examine the results is a simple graphical exploitation of the data. A resulting funnel plot reflects the distribution in Figure 1, which reports the absolute value of inequality of self-reported health studies plotted against a precision measure, which is the inverse of the standard error of the regression. Studies with less precision and hence, larger standard errors,
are at the bottom of the graph and will produce estimates that are more spread out. **Figure 1** makes apparent that there are large differences in the precision of inequality estimates, ranging from 0.2 to 0. Furthermore, it appears as though there were two distributions in the analysis that superimpose each other, one with a concentration index that is very close to zero and another distribution centred on 0.1. However, from simply observing a Funnel plot, it is not possible to ascertain the nature of such a distribution. The latter paves the way to pursuing a meta-regression strategy to investigate the underlying difference in inequality estimates. MRA will allow us to control for potential variables that explain the distribution of average inequality estimates. Finally, the lack of symmetry around the true coefficient (as shown by the plot with two funnels) is an early indication of publication bias.

**Figure 1. Funnel Plot (CI on X – axis and 1/SE on Y -axis)**

![Funnel Plot](image)

Source: own elaboration for study estimates

As the second step, we ran several meta-regression specifications, and performed the conventional FAT–PET tests as in equations (6), which are reported in Table 2. These tests will allow us to identify early the presence of publication bias and whether robustness of the empirical estimates is an issue.

Results from Table 2 suggest that the coefficient of the intercept is significant and suggests that we can reject the null hypothesis of no publication bias. However, estimates differ depending on whether regression estimates have clustered the standard errors by belonging to the same study, alongside a battery of controls. The significance of the intercept coefficient suggests that irrespective of the controls we adjust the mean inequality estimate for; there is still evidence of publication bias.
Controls include the way in which health system is financed (ie. whether estimates refer to an NHS country that does not exhibit a significant coefficient), the year of data of each estimate (suggesting the presence of inequalities increasing over time), the number of observations (which importantly does not seem to influence the regression results), whether the health data was cardinally measured (which appears consistently significant), and finally, whether or not the data has both a panel format (which does not appear significant).

The coefficient for 1/SE reflects the precision of the MRA or the so–called PET (precision-effect test), suggesting that the concentration index ranges from 0.016 to 0 depending on the controls and the clustering of the standard errors. Unfortunately, given the downward bias when there is a genuine effect (Stanley, 2008), these estimates calls for further testing. From results, we identify that the variable measuring the extent to which self-reported health was measured on a cardinal scale appears as significant in the specifications reported in Table 2. These results are indicative that possibly some source of bias lies in how health is cardinalised, when it is cardinalised. Furthermore, the significance of the intercept suggests that we can reject the null hypothesis of no selection bias (according to the FAT - funnel-asymmetry test) even when more controls are taken into consideration.

### Table 2. Funnel Asymmetry Test (FAT) and Precision Effect Test (PET)

<table>
<thead>
<tr>
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<th>coeff</th>
<th>Coeff</th>
<th>Coeff</th>
</tr>
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<tr>
<td></td>
<td>(s.e)</td>
<td>(s.e)</td>
<td>(s.e)</td>
</tr>
<tr>
<td>I/SE</td>
<td>0.013*</td>
<td>0.007</td>
<td>0.006</td>
</tr>
<tr>
<td></td>
<td>(0.005)</td>
<td>(0.005)</td>
<td>(0.005)</td>
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<tr>
<td>NHS</td>
<td>-1.849</td>
<td>-1.641</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(1.491)</td>
<td>(1.154)</td>
<td></td>
</tr>
<tr>
<td>Year of data</td>
<td>0.0379</td>
<td>0.008</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(0.0383)</td>
<td>(0.060)</td>
<td></td>
</tr>
<tr>
<td>N</td>
<td>-0.00011</td>
<td>1.17E-05</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(9.27E-4)</td>
<td>(6.74E-05)</td>
<td></td>
</tr>
<tr>
<td>Cardinal</td>
<td>-5.624*</td>
<td>-6.823*</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(1.453)</td>
<td>(1.369)</td>
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<tr>
<td>Panel</td>
<td></td>
<td>2.651</td>
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<tr>
<td>Intercept</td>
<td>2.155*</td>
<td>8.1547*</td>
<td>7.596*</td>
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<tr>
<td></td>
<td>(1.13)</td>
<td>(2.888)</td>
<td>(2.542)</td>
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<tr>
<td>Study cluster</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>F- Test</td>
<td>6.55</td>
<td>17</td>
<td>194.2</td>
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<tr>
<td>Adjusted $R^2$</td>
<td>0.15</td>
<td>0.47</td>
<td>0.52</td>
</tr>
</tbody>
</table>

* Highlighted if significant at least at 5%.

Notes: 1/SE refers to a measure of precision of the inequality estimate reported in each study. NHS refers to a dummy variable taking the value of 1 if the estimate refers to a health system financed by taxes. Year of data refers to the year the estimate refers to. Cardinal refers to a dummy variable to account for the cardinalisation of an inequality estimate. Finally, Panel refers to a dummy variable to measure whether the estimate has been computed using longitudinal data, and hence whether it filters potential unobserved heterogeneity.

In order to further filter the inequality indices for potential precision effects, Table 3 provides the estimates of the so-called precision effect estimate with standard error model (obtained as in equation 7). The coefficient for precision effects (1/SE) refers to the precision-corrected concentration index coefficient; that is, the concentration index corrected by selection bias, which lies between 0.013 and 0.0 depending on the specific study controls that are introduced. However, the most important effect we capture refers to the corrected concentration index after standard error clustering, suggesting that study-specific variability is more important than study characteristics such as the number of observations and other. One potential explanation of such results is the different degree of precision of different estimates, given that they rely on different samples and empirical strategies. Another important feature is that the inclusion of controls is increasing (decreasing) the value of the intercept (inverse of SE), suggestive of potential sources of unobserved heterogeneity in the measurement of health inequalities.
Table 3. Precision Effect Estimate with Standard Error (PEESE)

Note: * Significant at least at 5%.
Notes: 1/SE refers to a measure of precision of the inequality estimate reported in each study. NHS refers to a dummy variable taking the value of 1 if the estimate refers to a health system financed by taxes. Year of data refers to the year the estimate refers to. Cardinal refers to a dummy variable to account for the cardinalisation of an inequality estimate. Finally, Panel refers to a dummy variable to measure whether the estimate has been computed using longitudinal data, and hence whether it filters potential unobserved heterogeneity.

<table>
<thead>
<tr>
<th></th>
<th>coeff</th>
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<th>coeff</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>(s.e)</td>
<td>(s.e)</td>
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</tr>
<tr>
<td>SE</td>
<td>0.013*</td>
<td>0.013*</td>
<td>0.007*</td>
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<td>(0.004)</td>
<td>(0.006)</td>
<td>(0.002)</td>
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<tr>
<td>nhs</td>
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<tr>
<td></td>
<td>(0.54)</td>
<td>(1.124)</td>
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<td>0.0145</td>
<td></td>
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<td></td>
<td>(0.101)</td>
<td>(0.062)</td>
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<tr>
<td>N</td>
<td>1.59E-05</td>
<td>1.59E-05</td>
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<tr>
<td></td>
<td>(6.4E-05)</td>
<td>(6.67E-05)</td>
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<tr>
<td>Cardinal</td>
<td>-6.76*</td>
<td>-6.767*</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>(0.92)</td>
<td>(1.383)</td>
<td></td>
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<tr>
<td>Panel</td>
<td>2.694*</td>
<td>2.693</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(0.764)</td>
<td>(2.361)</td>
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</tr>
<tr>
<td>I/SE</td>
<td>1.837*</td>
<td>1.837</td>
<td>6.835*</td>
<td>6.834</td>
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<tr>
<td></td>
<td>(0.658)</td>
<td>(1.985)</td>
<td>(1.801)</td>
<td>(2.651)</td>
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<td>Study cluster</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>F- Test</td>
<td>4.31</td>
<td>7.99</td>
<td>178.2</td>
<td>234.7</td>
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<tr>
<td>Adjusted R²</td>
<td>0.15</td>
<td>0.3</td>
<td>0.42</td>
<td>0.52</td>
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</table>

dummy variable taking the value of 1 if the estimate refers to a health system financed by taxes. Year of data refers to the year the estimate refers to. Cardinal refers to a dummy variable to account for the cardinalisation of an inequality estimate. Finally, Panel refers to a dummy variable to measure whether the estimate has been computed using longitudinal data, and hence whether it filters potential unobserved heterogeneity.

4. Conclusions

This paper is to the best of our knowledge the first attempt to estimate the extent of publication bias of self-reported concentration index estimates. This is an important endeavour because there is some debate especially if we attempt to compare inequality estimates from different studies, given that study characteristics, system specific controls and robustness effects need to be controlled for to make inequality comparisons meaningful. In addition, the concentration index as a measure of health inequality is problematic insofar as it is a rank dependent measure and hence, presence of bias might in the estimates might be an
additional reason to rethink its use, and instead consider alternative measures (Costa-Font and Cowell, 2013).

The results of the study are important given that there is no clear view on what is the current level of inequalities in health in European countries: alternative cross-country analyses provide different results. One might expect heterogeneity in inequality estimates to result from study and empirical methodologies followed country specific effects, as well as the reliance on different health variables, heterogeneous databases and health system specific designs. If measures of inequalities in self-reported health reported in the literature were not corrected for methodological differences, comparisons of these measures across countries would not be appropriate, given that the data/methods used to obtain inequalities in health for each country will imply different types of measurement errors. The existing high heterogeneity and measurement error in the estimates shown in the literature on socioeconomic inequalities in health can be an issue in undertaking cross country comparisons, and potentially to estimate the effect of public policies on health inequalities.

This paper draws upon meta-regression analysis (MRA) to examine the influence of publication bias alongside precision and other study specific effects on estimates of income-related health inequalities. We rely on a sample of concentration index estimates and self-reported health measures, which is the common practice in the health economics literature. Our findings suggest evidence of publication bias that primarily depends on the cardinalisation of self-reported health. The latter is important because rank dependent inequality indices such as the concentration index are not meaningful unless indexes are cardinalised or turned into a ratio scale (Erreygers, T. van Ourti, 2011).

We find an effect from study-specific precision. We take advantage of an existing peer-reviewed literature on estimates of inequalities in health for different countries in Europe but these estimates have not been corrected and hence, comparisons across studies cannot be performed as they have different characteristics (including: year of the study, journal of publication, health variable used, inequality value, precision (standard error) of the estimated level of inequalities in health, among other factors). To date, there has been no analysis of this potential publication bias and subsequent correction of the measure of socioeconomic inequalities in health. By applying appropriate statistical methods, we are able to provide more comparable estimates of inequalities in health for each country. Once these corrected
measures are provided, it is possible to make more valid comparisons of the ranking of countries according to the adjusted measures of health inequalities. It may also be possible to identify publication and other biases in research on health inequalities.

We organise the literature by creating a database with all cross-country studies that provide estimates of socioeconomic inequalities in health, including details such as: the estimated level of inequalities in health, the precision of this estimate (standard error), the year of the study’s publication, the journal, the health variable used, the country analysed, the sample size used and several variables that will identify how those inequality measures were obtained. This information is analysed using meta-regression analysis (MRA). MRA entails a regression analysis of existing studies of socioeconomic inequalities in health, where the control variables are the type of study, the sample characteristics and the scope and precision of the estimate of socioeconomic inequalities in health, among others. MRA allows us to test the sensitivity of the estimate of inequalities in health to the study characteristics\(^5\).

MRA is especially designed to allow correcting empirical estimates, in our case, measures of socioeconomic inequalities in self-reported health for potential biases. By creating a uniform structure for scrutinizing studies, our work attempts to make an important contribution to the literature on inequalities in health. Correcting for publication biases appear as particularly relevant when inequality estimates are employed in ranking health systems or simply when comparing estimates across countries, an issue that will be of interest to policymakers. Furthermore, once a corrected measure of inequalities in health has been attained, one can used such corrected estimates to contribute to research debates, such as those on the equity-efficiency trade-off, by providing corrected inequality values that can be used in any analysis.

Finally, this study suggests important avenues for policy in so far as it indicates that there is some consensus in the methods employed. The latter might well be the results is using similar self-reported measures of health status and common measures of income. However, the existence of self-reporting bias in both health and income measurement still remain an issue (Costa-Font and Hernandez-Quevedo, 2013). Overall results are consistent with

\(^5\) No doubt, there are other important factors affecting the MRA coefficients which are worth to mention and that should be accommodated in the estimations (i.e. omitted variable bias, unobserved heterogeneity, functional form, etc.)
evidence of persistence inequalities in health care which suggest the need for path breaking intervention designs.

References


APPENDIX

**Table A1. List of Studies Included in the MRA**


