Dear Dr. Jens-Arne Subke,

We are pleased to submit our revised manuscript, following your and the referees’ suggestions, along with a marked up copy in which all changes since the original submission are detailed in red. Below you will also find our point-by-point responses to your comments and to the referees’ comments (note that you have already read our responses to the referees’ comments; we include them again here only because the instructions for this stage say we should). We have not included a separate list of all changes because it would be redundant with (and surely less useful than) the marked up manuscript.

We thank you for your efforts on our manuscript. Please let us know if you would like anything else from us.

Best Regards,

Rick Wehr (also on behalf of Scott Saleska)

Response to the Associate Editor’s Decision on bg-2016-315

Response of authors Wehr and Saleska (hereafter “WS) to Associate Editor Jens-Arne Subke (hereafter “JS”):

JS: Many thanks for your replies to the two referee reports. Both referees have made an excellent job of appraising your manuscript critically, and together with your response to their individual points raised, I think it has made this publication even stronger. I’m sure that in its final form, it will become an important reference for isotope biogeochemists.

WS: Thank you. We agree that the referee’s and your comments have improved the manuscript, and we hope that it will be well received by readers.

JS: I’d like you to implement the suggested revision points, including a brief discussion of the issue of instrument uncertainty vs. variability caused by spatial/ecosystem heterogeneity, raised by referee 2 (John Miller).

WS: We have implemented all the suggested revision points in our revised manuscript, including the addition of a new section discussing natural variability (Section 2.2).

JS: The new figure illustrating a ‘real data’ example of night-time Keeling plots is also very useful. A detailed figure caption outlining the differences in data sets would obviously be necessary alongside it.
We have included the new figure in our revised manuscript, with a caption and a corresponding discussion in the text (Section 4.2).

Response to Referee Comments on bg-2016-315 (already seen by the Associate Editor)

Author Response to Anonymous Referee #1 [hereafter “AR1”]

AR1: The manuscript by Wehr and Saleska re-introduces a non-linear iterative method to determine slope and intercepts of mixing line relationships. Isotopic mixing line relationships have been analyzed in previous studies (notably Zobitz et al. 2006 and Kayler et al. 2010). The current study expands on the previous two by introducing a “long-solved” method. Overall the paper is well written and readable. A strength of the manuscript is that it very acutely emphasizes the disconnect between the geoscience and environmental science - and arguably the mathematical science - communities. Technical advances in one area don’t seem to percolate over the the other area (as highlighted in the second paragraph of the introduction). Illuminating this tension between translatability across disciplines is a strength of the paper.

We thank the referee for these supportive comments, and for taking the time to review our manuscript.

AR1: There are few weaknesses to the paper which could be addressed by revision.

We appreciate the referee’s suggestions and will incorporate them into our revised manuscript as described below.

AR1: First, there is a lack of recognition of the importance of OLS and other regression methods. OLS, GMR, ODR, as well as Maximum Likelihood Estimation are essentially a linear problem and are amenable to several different approaches in mathematics - OLS is a topic in a Calculus sequence. The York method, best I can tell, is a non-linear iterative method - which perhaps may contribute to its unfamiliarity across disciplines.

We agree on “the importance of OLS and other regression methods”, and had intended that the manuscript acknowledge their wide use while making clear that their overuse (in cases where their assumptions do not apply) is an explicit motivation for the paper. Regarding the contrast between methods, yes, the general LSE solution for the slope and intercept cannot be written analytically, and so York’s method finds it numerically, by iteration. The OLS, GMR, and ODR slopes and intercepts are special cases of the general LSE solution that can be solved for analytically. We can mention this distinction in our revised manuscript.

AR1: Second, I also think some more careful tracking of the timing of the key studies cited is important. Zobitz et al 2006 was written in response to previous studies by Pataki et al 2003. Kayler et al 2010 addressed minimizing bias for large CO2 ranges -
and addressed some of the issues raised in the previous two studies. A consistent finding both in Zobitz et al 2006 and Kayler et al 2010 is that OLS is appropriate for sufficiently large CO2 ranges and is highly biased at low CO2 ranges. Given that, the authors of the current manuscript don’t present a pressing need to move away from OLS in favor of a more complicated linear fitting method. What is the current state of the art in the measurement method? How imperative to determine mixing line parameters with samples of low CO2 ranges? Addressing some of the importance and need of this method will help increases its applicability, and the tradeoff for using a relatively more complex fitting routine than what is provided on all statistical software programs (R, SAS, etc).

We like the referee’s suggestion to add mention of the timing/context/motivation of the cited studies, and we will do so in our revised manuscript. Zobitz et al 2006 actually reported that OLS was negligibly biased for all measurement conditions (but the random error increased at low CO2 ranges). On the other hand, Kayler et al 2010 reported that the OLS Keeling plot intercept was negligibly biased (i.e. by less than about 0.1 permil) for CO2 ranges above roughly 50 ppm but non-negligibly biased for CO2 ranges below roughly 10 ppm. The particular numbers depend on the measurement precisions (as illustrated by our tables); however, in Kayler et al 2010, various precision scenarios seem to have been amalgamated in their Table 1, so that for a given set of measurement precisions, it is not clear whether non-negligible OLS bias emerges at a CO2 range of 100 ppm or 10 ppm. That is an important issue because Keeling plots with CO2 ranges below 50 ppm are not unusual — indeed the real data example that we will include in our revised manuscript (see below) involves hundreds of Keeling plots made from nighttime air sampling in a forest, and most of those Keeling plots have a CO2 range below 50 ppm (for quite a few, the range is even below 25 ppm). So prior to the present manuscript, it was unclear whether it was acceptable to use OLS under some fairly common conditions. Moreover, the primary conclusion of Kayler et al 2010 was actually that GMR is better than OLS for very high CO2 ranges such as might be encountered in soil measurements (“The combination of geometric mean regression and the Miller–Tans mixing model provided the most accurate and precise estimate of d13CR when the range of CO2 is >1,000 umol mol-1.”). The main goal of our paper is to inform readers that there is a single general fit method (York’s) that is best in all measurement scenarios, so that they do not need to make a choice among imperfect methods or to switch between those methods depending on the conditions. We will add a paragraph to the introduction clarifying the above points, thereby adding motivation for the use of York’s method.

Having said all that, we recognize that OLS is more accessible and familiar to researchers, and we do not argue that researchers must abandon OLS. Indeed, a second goal of our paper is to accurately detail the range of conditions in which OLS can safely be used to approximate the York solution (hence the tables with biases under various conditions). It turns out that under most practical conditions, OLS is an acceptable approximation to the York solution, while under some conditions, it is not. We will clarify this point in the Results and discussion section. When in doubt,
researchers will now have a general method that they can use without risking introducing bias.

More broadly, the isotopic mixing line is being used here as a practical example to illustrate the York method, which might prove even more valuable to other applications, quite apart from our own line of research. We would like to add a paragraph briefly presenting a separate potential application of York’s solution, which is quite common across disciplines: the comparison of measurements of the same quantity by two different instruments that differ in their precisions (such as when replacing an old instrument with a new one).

**AR1:** Third, the results of this paper relied on subsampling of simulated data, which does limit the applicability of their results. I suggest the authors provide a case study of non-simulated data, comparing the two York method to OLS, GMR, ODR etc. Simulations are great for emphasizing the theoretical underpinnings of a method, however the addition of real measured data would enhance the applicability and impact of the simulation results.

As mentioned above, we have a real data set that we will add to the manuscript: precise (0.05 permil and 0.02 ppm) nighttime air sampling on the tower at the Harvard Forest Environmental Measurements Site. Given that our simulated data results show the York method to be unbiased, we can suppose that the bias in GMR (or OLS) is given by the difference between the GMR (or OLS) intercept and the York intercept. The 429 nights in our dataset (with one Keeling plot per night) allow us to clearly see the dependence of the biases on the CO2 range — and by adding additional noise to the real data, we also get an indication of the dependence of the bias on the instrument precision. We find that OLS is a very good approximation to the York solution for our particular instrument precision, while GMR is a poor one. With noise levels typical of lower-cost instruments, we find that even OLS becomes a poor approximation for CO2 ranges less than 10 ppm, in agreement with our simulated data results. These results using real data are shown in the figure attached to this response.

**AR1:** P1 L21: “Much of it was outdated before it was written” is a very vague sentence.

Taken in isolation, it is vague, but this sentence is the deliberately suspenseful setup for the two sentences that follow, which make the point that the solution sought by the literature in question was already known in 1969. In context, we feel that the sentence is not vague, and that it adds to the readability of the manuscript.

**P1 L26:** Point made that York’s solution is unknown, but impact is not an indication of quality - I think it just got dwarfed, highlights the need for interdisciplinary collaboration.

We are glad that the referee agrees with us on this motivational point.
AR1: P2 L9: Delete "but the debate is immaterial" This is a general sentence that is unprovable and opinion.

We meant to convey the idea that when a convenient, general solution is available, it is not necessary to choose among special-case solutions. But perhaps we do not need to make that point here, and so we will delete this phrase.

AR1: P3 L1: Please clarify if the Hirsch and Gilroy citation applies to all the quoted phrases in this sentence or only one (clarify)

We will clarify by writing: “Adding confusion to the literature is the fact that OLS, ODR, and GMR are each known by other names (York, 1966; Hirsch and Gilroy, 1984): OLS is called….”.

AR1: P4 L27: Given the fact you need an initial guess slope, is the convergence of the method sensitive to the initial guess value, or does it converge globally?

The method will require more iterations to converge to the desired accuracy level when the initial guess is poorer, but the value of the initial guess has little impact on the convergence speed unless the guess is extreme. As stated in the manuscript, “This guess can be very rough and still sufficient... If desired, a good initial guess slope can be obtained from an OLS fit”.

AR1: P7 L24: “For CO2 ranges less than 50 ppm…” This sentence reads very awkwardly and to follow the logic.. Please rephrase

We can rephrase the sentence thusly: “For CO2 ranges less than 50 ppm, the GMR bias is non-negligible (> 0.1 ‰) unless the measurement uncertainty in δ is extraordinarily low (≤ 0.01 ‰).”

AR1: P8 L22: Now I am confused. Does the York method give an exact solution (as in OLS, GMR) or is it a nonlinear iterative method as described on page 4?

We see that the meaning of ‘exact’ was unclear. York’s general LSE solution is exact in that no approximation is involved in its derivation, unlike the preceding approach of Deming (1943), which dropped terms from a Taylor expansion. As mentioned above, York’s solution is also a numerical, iterative solution, in contrast to the special-case LSE solutions (OLS, ODR, and GMR), which can be written analytically (i.e. in closed form). In practice, there is no difference in exactitude between the numerical approach of York and the analytical approach of OLS, ODR, and GMR; all are limited only by machine precision. This will be clarified in the revision.

Author Response to John Miller (Referee) [hereafter “JM”]
JM: Congratulations on writing a terrific paper: well researched, well written and very much needed. I have just a few questions and comments.

We thank the referee for his compliments and for taking the time to review our paper.

JM: How can we, or should we, consider variability in CO2 and δ13C arising not from instrumental noise but from the environment? As pointed out in Miller and Tans, in some real-world situations, assignment of analytical uncertainties to CO2 and δ13C may result in poor goodness of fit, i.e. a large value of reduced-chi square, suggesting that analytical CO2 and δ13C uncertainties are too small. This is of course important because small CO2 and δ13C uncertainties will lead to too small slope and intercept uncertainties. Note that while not so common now, as analytical precision improves, instances where natural variability significantly exceeds instrumental precision will need to be dealt with more. In MT2003, we attempted to deal with this by starting with an initial estimate of the best fit line, although we used a GMR instead of fitexy (for speed, and because we only knew the analytical uncertainties). We then proceeded to scale the standard deviations of the x and y residuals to produce a reduced chi-square value of 1; finally fitexy was used to calculate the slope and intercept uncertainties. Nonetheless, a problem persists, which is that the slope of the best fit line depends on the initially assigned x and y uncertainties. I’m very interested to hear your ideas of how to address this. (Maybe I’m missing something obvious, like using an OLR regression as a starting point.)

As you know, York’s method deals with the situation in which there is a linear relationship between the true values of X and Y (i.e. of CO2 and d13C) but those values are measured with error. Natural variability may sometimes be describable as measurement error; that is, as a stochastic process that intervenes between the quantity of interest and the measurement of that quantity. For example, the eddy covariance method uses a single-point measurement to estimate the gas flux through a large 2D plane overlying an ecosystem, and most of the noise in the estimation comes not from the instrumental measurement uncertainty but from the natural (turbulence-driven) variability in the flux past that single point relative to the flux through the whole 2D plane. If the natural variability in X and the natural variability in Y are describable as measurement error and can be characterized independently (along with any correlation between them), then York’s method can be applied and is likely to be very useful.

On the other hand, it is often the case that the natural variability is not well characterized, or that it is not well described as additional measurement error. In this case, we argue that one cannot proceed to determine the best-fit line, or even to define what “best-fit” means. In general, one can view natural variability as variation in the true X-Y relationship due to the influence of other factors that are not controlled for. So a Keeling plot with natural variability is like many true mixing lines all superimposed on the same plot (one line for each set of influencing factors).
It is therefore pertinent to consider which true line one is looking for. To define that line is, in effect, to characterize the natural variability in $X$ and $Y$.

If one is interested not in the $X$-$Y$ relationship per se (i.e. not in an intercept or slope), but simply in predicting the most likely value of $Y$ given $X$ for the particular data that were sampled and put on the plot, then OLS is the proper fit to use. If differences among the various fit methods are not large enough to matter to the scientific question being posed, then OLS is again a sensible choice, owing to its simplicity.

If it would be helpful for context, we would be happy to include a brief discussion of the above points in the manuscript.

Regarding the approach you mention from MT2003, it seems that the slope you end up with must depend on your arbitrary choice of how to apportion the variability into $X$ and $Y$. That seems to be what you are saying when you say that "the slope of the best fit line depends on the initially assigned $x$ and $y$ uncertainties". Any meaningful approach will require independent information on how to apportion the natural variability between $X$ and $Y$.

\textit{JM: P4 l8. Note for future reference that the Keeling plot equation is valid not just for a single source (or sink), but \textit{delta}_s can be interpreted as a flux-weighted source (sink) signature.}

That is a good point. We will remove the word “single” from this sentence.

\textit{JM: Eq. 3. The derivation of this was not obvious. It’s not critical to the argument, but since you have an appendix, can you add this?}

Yes, we will be happy to add the derivation of Eq. 3. It consists merely of writing the definition of the correlation coefficient in terms of the covariance, and then the definition of the covariance in terms of means and expectations, and so it is probably short enough to be included in the main text.

\textit{JM: P7 l5. I am surprised by (and skeptical of) an instrument with 0.01 ppm and 0.01 per mil uncertainty. Can you provide a reference in the literature for this, especially since this is characterized as ‘common instrumentation’?}

The 0.01 ppm and 0.01 permil uncertainties are attributed to "the best existing laser spectrometer" rather than common instrumentation, but we recognize that this point is confusing because of the placement of the phrase “including some corresponding to common instrumentation" (which was meant to apply to the whole list). We will move that phrase to avoid the confusion. The spectrometer referred to has a precision of 0.016 ppm and 0.02 permil under optimal conditions, and we will add its citation (Wehr et al, 2013, Agricultural and Forest Meteorology, 181,69–84). We will also clarify that 0.01 and 0.01 are slightly better than said
spectrometer (we chose to round down to 0.01 because our aim was to bracket the range of conditions that researchers are likely to encounter, and spectrometer precision is likely to continue to improve).

**JM: P7l8. Change ‘latter’ to ‘last’.

Ok, we will make this change.

**JM: P7l34 and Table 2. I’m confused as to why CO2 ranges from 100 to 5000 are relevant and why CO2 uncertainties greater than 1 are relevant. I understand that soil chambers could give such high CO2 enhancements, but as seen from the table, uncertainties become very small. Perhaps you could add a column of 100 ppm in Table 1 and then summarize the rest of the Table2 results in the text.

We included those ranges and uncertainties for comparison to Kayler et al 2010, where it is argued that high CO2 values are often accompanied by high uncertainties. We agree that the 5000 ppm column is unnecessary, but for the highest CO2 uncertainty (20 ppm), there are non-negligible biases even for a CO2 range of 1000 ppm. We can condense the table substantially by eliminating the 5000 ppm column and then putting the Keeling and Miller/Tans results side by side rather than on top of one another.

**JM: P8. L1. Why are the MT results a bit better at these high values? Or maybe better to say, why are the KP biases occasionally significant?

This question is the subject of the following paragraph in the manuscript (beginning on P8 L4).

**JM: P8l29. Factor of 2 seems a bit too generous. The biggest offset from Monte Carlo I see is 0.67.

We can say that “the agreement is nonetheless within 33%.”

**JM: P8l35. What are the ‘adjusted data points’?

The adjusted data points are the fit method’s estimation of the ‘true’ data points that were measured with error in order to produce the measured data points. We will add this explanation to the manuscript.

**JM: P9.17 Isn’t G simply reduced chi-square? If so, why introduce a new term for this?

G is the weighted reduced chi-square. In our revised manuscript, we will state that this is the goodness of fit metric being used, and we will use the variable chi-square in place of G.
The long-solved problem of the best-fit straight line: Application to isotopic mixing lines

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Abstract. It has been almost 50 years since York published an exact and general solution for the best-fit straight line to independent points with normally distributed errors in both x and y. York’s solution is highly cited in the geophysical literature but almost unknown outside of it, so that there has been no ebb in the tide of books and papers wrestling with the problem. Much of the post-1969 literature on straight-line fitting has sown confusion not merely by its content but by its very existence. The optimal least-squares fit is already known; the problem is already solved. Here we introduce the non-specialist reader to York’s solution, and demonstrate its application in the interesting case of the isotopic mixing line, an analytical tool widely used to determine the isotopic signature of trace gas sources for the study of biogeochemical cycles. The most commonly known linear regression methods—ordinary least squares regression (OLS), geometric mean regression (GMR), and orthogonal distance regression (ODR)—have each been recommended as the best method for fitting isotopic mixing lines. In fact, OLS, GMR, and ODR are all special cases of York’s solution that are valid only under particular measurement conditions, and those conditions do not hold in general for isotopic mixing lines. Using Monte Carlo simulations, we quantify the biases in OLS, GMR, and ODR under various conditions and show that York’s general—and convenient—solution is always the least biased.

1 Introduction

A common analytical task in the physical sciences is to find the true straight-line relationship underlying independently measured points with normally distributed measurement errors in both the ordinate y and abscissa x. The literature on this topic is profuse, but much of it was outdated even before it was written. Looking through it, one discovers a menagerie of least-squares regression methods, none strictly appropriate to the task at hand. What one is unlikely to find, unfortunately, is the general, exact, and convenient least-squares solution published in 1969 by geophysicist Derek York (York, 1969).

York was motivated by rubidium-strontium isochrons but his landmark solution was universal. Unfortunately, while it became standard in geophysics, York’s solution has remained largely unknown to the broader scientific community: York’s paper has been cited almost 2000 times within the geophysical literature and only about two dozen times in all the rest of the scientific literature combined. Meanwhile, the number of papers and chapters expounding on the subject as if the solution did not exist is staggering. Examples include widely consulted books like Biometry (Sokal and Rohlf, 1995) and Numerical Recipes (Press et al., 2007) as well as articles in fields as diverse as: anthropology (Smith, 2009), water resources (Hirsch and Gilroy, 1984), clinical chemistry (Stöckl et al., 1998), marine biology (Law and Archie, 1981), aerosol science (Leng et al., 2007), and astronomy (Feigelson and Babu, 1992).

One scientific problem wanting for York’s solution is the isotopic mixing line. In 1958, Charles Keeling (Keeling, 1958) introduced the idea of using an isotopic mixing line to determine the stable isotopic signature of a trace gas source. If the isotopic
composition $\delta$ of a trace gas with a net source or sink is plotted versus the inverse of its mixing ratio in air $c$, the points describe a straight line whose $y$-intercept gives the desired source or sink signature. The “Keeling plot” became a key method in isotope ecology and biogeochemistry, and is regularly used, for example, to determine the isotopic composition of respired CO$_2$. Unaware of York’s general least-squares solution, researchers concerned with Keeling plots have debated which of the more widely known, special-case least-squares regression methods is best. Some studies have concluded in favor of ordinary least squares regression (OLS) (Zobitz et al., 2006; Kayler et al., 2010), others in favor of orthogonal distance regression (ODR) (Ogée et al., 2003; Miller & Tans, 2003), and still others in favor of geometric mean regression (GMR) (Pataki et al., 2003; Miller & Tans, 2003; Kayler et al., 2010). The disagreement arises because in each of these special cases, the measurement conditions influence the fit line bias.

Here we introduce the reader to York’s solution and its practical application, using the isotopic mixing line as a concrete example. We then use Monte Carlo simulations to precisely quantify the biases inherent to York’s solution and to the popular special-case regression methods under various common (and some uncommon) conditions. In Appendix A, we provide a short, fast, and easily implemented algorithm for computing York’s best-fit slope and intercept, as well as their associated uncertainties.

2 Background

2.1 The taxonomy of straight-line fitting methods

The goal of straight-line fitting is to retrieve the true straight-line relationship between two variables, $x$ and $y$, from their measured values, $\hat{x}$ and $\hat{y}$ (hats will denote measured values throughout this paper). Though it is common to neglect the error in $\hat{x}$, doing so can lead to a biased fit, as both $\hat{x}$ and $\hat{y}$ are measurements and therefore corrupted by error at some level. The nature of that error might vary from point to point (a situation known as heteroscedasticity), and it might also be that the error in $\hat{x}_i$ is correlated with that in $\hat{y}_i$, where the subscript $i$ specifies a measurement pair (i.e. a data point). Such correlation can arise, for example, if both $\hat{x}$ and $\hat{y}$ are derived from the same quantity or measured by the same apparatus, and can be described by a (Pearson’s) correlation coefficient $r$ that might also vary from point to point. Finally, it is conceivable that the errors in the various $\hat{x}_i$ might be correlated with one another, and similarly for the $\hat{y}_i$; that is, the points might not be independent.

If the points are independent of one another and their errors are normally distributed, then the problem can be treated by Least-Squares Estimation (LSE), which is equivalent to Maximum Likelihood Estimation (MLE) (Myung, 2003) in this situation. Most of the literature on straight-line fitting concerns LSE, as it is appropriate to the vast majority of straight-line fitting problems.

York’s solution is the general LSE solution: his equations provide the best possible, unbiased estimates of the true intercept $a$ and slope $b$ in all cases where the points are independent and the errors are normally distributed. We refer to this general solution as exact because no approximation is involved in its derivation, although it cannot be written analytically and must therefore be obtained in practice by numerical iteration. In contrast, the LSE methods commonly used and debated—namely OLS, ODR, and GMR—provide unbiased estimates only in very specific special cases in which the solution can be written analytically. York (1966, 2004) considers those situations mathematically and shows that his equations reduce algebraically to OLS, GMR, etc. when appropriate.
Before discussing OLS, ODR, and GMR further, we should note that each is known by other names that add confusion to the literature (York, 1966; Hirsch and Gilroy, 1984): OLS is called ‘the regression of y on x’, ODR is called ‘major axis regression’, and GMR is called ‘reduced major axis regression’. ODR has also been called ‘least normal squares’ and ‘the line of closest fit’, while GMR has also been called ‘the line of organic correlation’, ‘the unique solution’, and ‘the equivalence line’. The methods are also often categorized, with methods that consider error only in y (e.g. OLS) being called ‘Model I’ regressions, and those that consider error in both x and y (e.g. ODR, GMR) being called ‘Model II’ regressions (Sokal and Rohlf, 1995).

OLS is by far the most widely known fitting method. The OLS fit line is unbiased only when there is negligible error in x and when the error variance for the \( y_i \) does not vary with \( i \). In this case, the problem reduces to minimizing the sum of the vertical distances of the points from the fit line. A variant known as weighted least-squares allows the error variance for the \( y_i \) to vary with \( i \).

The ODR fit line (Pearson, 1901) is unbiased only when the error variances for the \( x_i \) and the \( y_i \) are equal and independent of \( i \), and when the error in \( x_i \) is uncorrelated with that in \( y_i \). (The technique Pearson invented to arrive at the ODR fit line is called principal components analysis, which is widely used in its own right.) In this case, the problem reduces to minimizing the sum of the squares of the perpendicular distances of the points from the fit line. However, the condition of equal error variances is almost never satisfied in reality. The excessively restrictive nature of ODR is highlighted by the fact that the ODR result is not invariant under a change of scale. In other words, if one scales the y-axis by a factor of 10, the fit line slope does not scale by a factor of 10. This flaw led to the development of GMR (Kermack and Haldane, 1950), which effectively performs ODR on transformed coordinates: the \( x_i \) divided by the standard deviation of the \( x_i \), and the \( y_i \) divided by the standard deviation of the \( y_i \) (not by the standard deviations of their errors). However, GMR is also highly restrictive, as its fit line is unbiased only in the peculiar circumstance that the variance of the \( x \) error divided by the variance of the \( x_i \) is equal to the variance of the \( y \) error divided by the variance of the \( y_i \).

Neither OLS, ODR, nor GMR uses estimates of the actual measurement uncertainty in its determination of the best-fit line.

A superior fitting method, called fitexy, is provided as an algorithm in Press et al. (2007) and its earlier editions. Press et al. (2007) started (unknowingly) from a similar point as did York (1969), seeking to minimize a \( \chi^2 \) function that is identical to \( S \) in Eq. (2) of York (1969) if the correlation coefficient there is set to zero. However, rather than taking advantage of York’s (1966, 1969) algebra, Press et al. (2007) treat the problem as one of nonlinear minimization. This approach leads them to a more complicated (and slower) algorithm than one based on York’s solution (such as that provided here in Appendix A). Moreover, fitexy is less general than York’s solution in that it does not allow the errors in \( x_i \) and \( y_i \) to be correlated. The fitexy method has sometimes been confused with ODR in the literature (Ogée et al., 2003; Pataki et al., 2003).

If the errors in any or all of the \( x_i \) and \( y_i \) are known to be distributed non-normally, then strictly speaking, LSE does not apply and one should retreat to the even more general formalism of MLE. However, the bias introduced into York’s solution by non-normal error distributions is not always a problem, as we show in Section 4.1. The bias will vary on a case-by-case basis, but if the correct distribution is known, then the significance of the bias can be estimated fairly simply by Monte Carlo simulation, as we do here.
If LSE must be rejected, MLE may or may not be tractable. MLE requires that the correct error distributions be written analytically, and that a useable expression be derived for the likelihood function $L$ appropriate to those distributions. We say ‘useable’ because simply applying the definition of $L$ to the case of a straight line with errors in both $x$ and $y$ yields an expression that requires knowledge of the true values $x_i$ and is therefore useless in and of itself. Expressing the likelihood function in a form that does not include the true values $x_i$ was essentially York’s first step in deriving his solution (see Eqs. (7) through (14) of York (1966), in which $S \propto -\ln(L)$).

### 2.2 A note on natural variability

York’s method deals with the situation in which there is a linear relationship between the true values $x$ and $y$ but those values are measured with random error. When the scatter of the measured values $\bar{x}$ and $\bar{y}$ about a straight line exceeds the uncertainty attributed to the measurement technique, the excess scatter is attributed to “natural variability”. Natural variability may sometimes be describable as just another part of the measurement error; that is, as a stochastic process that intervenes between the quantity of interest and the measurement of that quantity. For example, a technique called eddy covariance is commonly used to estimate the flux of CO$_2$ through the two-dimensional plane overlying an ecosystem based on a single-point measurement on a tower, and most of the noise in the estimation comes not from the instrumental measurement uncertainty but from the natural (turbulence-driven) variability in the flux past that single point relative to the flux through the whole plane (Wyhr et al., 2013). If the natural variability in $\bar{x}$ and the natural variability in $\bar{y}$ are describable as normally distributed measurement error and can be characterized independently (along with any correlation between them), then York’s method can be applied and is likely to be very useful.

On the other hand, it is often the case that the natural variability is not well characterized, or that it is not well described as additional measurement error. In this case, we argue that one cannot proceed to determine the best-fit line, or even to define what “best-fit” means. In general, one can view natural variability as variation in the true relationship due to the influence of other factors that are not controlled for. So a Keeling plot with natural variability is like many true mixing lines all superimposed on the same plot (one line for each set of influencing factors). It is therefore pertinent to consider which true line one is looking for.

To define that line is, in effect, to characterize the natural variability in $x$ and $y$.

If one is interested not in the $x$-$y$ relationship per se (i.e. not in an intercept or slope), but simply in predicting the most likely value of $\bar{y}$ given $\bar{x}$ for the particular data that were sampled and put on the plot, then OLS is the proper fit to use. If differences among the various fit methods are not large enough to matter to the scientific question being posed, then OLS is again a sensible choice, owing to its simplicity.

### 2.3 The isotopic mixing line problem: Keeling and Miller/Tans plots

A Keeling plot is a scatterplot of the stable isotopic composition $\delta$ of a trace gas (typically CO$_2$) versus the inverse of its mixing ratio $c$ in air (Fig. 1). The isotopic composition $\delta$ is defined as a relative deviation from a reference material: $\delta = (R - R_{ref})/R_{ref}$, where $R$ is the ratio of the rare isotope (e.g. $^{13}$C) abundance to the most common isotope (e.g. $^{12}$C) abundance, and $R_{ref}$ is that ratio for the reference material. The standard units for $\delta$ are $\%$ (parts per thousand), while those for $c$ are ppm (parts per million).

It is easy to show that in the case of a source or sink changing the trace gas mixing ratio in the atmosphere, the $y$-intercept of a straight line fit to a Keeling plot gives the isotopic signature (i.e. the $\delta$) of that source or sink. To wit, if the subscripts $t$, 0, and $s$ refer to the atmosphere at some time $t$, the atmosphere at time 0, and the source, respectively, then:
$\epsilon_t = \epsilon_0 + \epsilon_i$
$\delta \epsilon_t = \delta \epsilon_0 + \delta \epsilon_i = \delta \epsilon_0 + \delta \epsilon_i (\epsilon_t - \epsilon_0)$
$\Rightarrow \delta_t = \delta_i + (\delta \epsilon_0 - \delta \epsilon_i) \times (1/\epsilon_i)$

Multiplying the Keeling fit line equation $\delta = a + b(1/c)$ through $c$ gives an alternate straight-line equation, $\delta c = b + ac$, which can be fit to a plot of $\delta$ versus $c$, called a Miller/Tans plot (Miller & Tans, 2003). In that case, it is the slope of the line that gives the source or sink signature. (Ironically, Keeling reported the inverse relationship between $\delta$ and $c$ in 1958 but no “Keeling plots” appear in any of his early papers, nor does he discuss straight line fitting per se; he chose instead to show curved fit lines to plots of $\delta$ versus $c$.)

In many ecosystems, the source/sink signatures of interest differ from one another by just 1‰ or less (Bowling et al., 2014), so that Keeling or Miller/Tans plot fit biases of 0.1‰ can be important.

The first studies to consider the effect of error in $c$ on isotopic mixing line fits (Miller & Tans, 2003; Pataki et al., 2003; Ogée et al., 2003) advocated use of ODR or GMR on the grounds that OLS is biased by neglecting error in $c$; however, they quantified only the differences between OLS and the other methods rather than their true biases. Zobitz et al. (2010) then examined the true biases through Monte Carlo simulations and reported that OLS was negligibly biased for all measurement conditions, and that the difference between OLS and GMR was in fact due to bias in GMR. Kayler et al. (2010) later revisited the issue and reported that OLS could be non-negligibly biased for both low (<50 ppm) and high (>1000 ppm) $CO_2$ ranges, depending on the measurement conditions, and they advocated use of GMR on a Miller/Tans plot for the most accurate fits when the $CO_2$ range is high. One goal of the present article is to inform readers that there is a single general fit method (York’s) that is best in all measurement scenarios, so that they do not need to make a choice among biased methods or to switch between those methods depending on the conditions. Another goal is to detail the conditions under which York’s general solution can be satisfactorily approximated by OLS, as we recognize that OLS will likely remain more accessible and familiar to most researchers.

3 Methods

3.1 Applying York’s solution to isotopic mixing lines

An algorithm that solves the York equations (see Appendix A) requires as input, for each data point $i$: the abscissa $x_i$, the ordinate $y_i$, the standard deviations of the errors for $x_i$ and $y_i$, and a correlation coefficient $r$, describing the correlation between the error in $x_i$ and that in $y_i$. This set of information is the same set required for any accurate straight-line fit—if a method does not ask for some of these parameters, then it is implicitly assuming values for them. However, as York’s equations must be solved by numerical iteration, they require one additional piece of information, an initial guess slope. This guess can be very rough and still sufficient, as convergence is not sensitive to the value chosen, and the solution can be iterated to an arbitrary degree of accuracy (using the previous best-fit slope as the new guess slope). In practice, ten iterations are almost always sufficient. If desired, a good initial guess slope can be obtained from an OLS fit.

The accuracy of the best-fit line will depend on the accuracy of the error and correlation estimates, but almost any reasonable estimates will be better than the estimates implicit in OLS, GMR, and ODR. (And good error estimates are to be sought anyway,
as a measurement is only meaningful to the extent that its uncertainty has been quantified.) The Keeling plot is an interesting application partly because, if we take the errors in $c$ to be normally distributed, then the errors in the Keeling abscissa $x = 1/c$ are not normally distributed—though the errors in $c$ tend to be small relative to $c$ itself, so that the resulting bias in the fit line tends to be negligible (see Section 4.1). Miller/Tans plots are also interesting, for a different reason: though we take the errors in $t$ to be normally distributed, the errors in the Keeling abscissa $x = c$ and ordinate $y = (dc)_t = c t$ are always correlated because both contain $c$. Moreover, the errors in the Keeling abscissa and Miller/Tans ordinate are both (slightly) heteroscedastic.

Regarding correlation between the errors in $x$ and $y$, it is perhaps worth mentioning that the high-frequency (e.g. 1 Hz) random noise in modern spectroscopic measurements of $\delta$ and $c$ is generally correlated due to the fact that certain causes of random spectroscopic noise (e.g. laser frequency instability) necessarily affect retrievals of both $\delta$ and $c$. The correlation may nonetheless be negligible for data averaged over a minute or longer.

The standard deviations of the errors in $c$ and $1/x$, as well as the Miller/Tans correlation coefficient, are given by Eqs. (1), (2), and (6), below. For a series of $i$ measurement pairs $(x_i, y_i)$, let the standard deviations of the normally distributed measurement errors be $(\xi, \eta_i)$. Then the standard deviation of the measurement errors in the Miller/Tans ordinate $(\delta c)$ is given by (again, assuming no correlation between the errors in $x$ and $y$):

$$
\phi = \sqrt{\xi^2 \delta_t^2 + \eta_i^2 \xi_i^2} \approx \sqrt{\xi_i^2 \delta_t^2 + \eta_i^2 \xi_i^2}.
$$

(1)

Meanwhile the standard deviation of the (not really normally distributed) measurement errors in the Keeling abscissa $(1/c)$ is given by:

$$
\theta = \xi_i/c_i^2 = \xi_i/c_i^2.
$$

(2)

Eqs. (1) and (2) are standard expressions for the propagation of uncertainty through the operations of multiplication and division. For a Keeling fit, the correlation coefficient $r_i$ should be set to zero for all $i$ (unless there is correlation between the measurement errors in $\delta$ and $c$). For a Miller/Tans fit, $r_i$ is defined as:

$$
r_i \equiv \text{cov}(\delta c, \xi) / \phi \xi_i = (\delta c) \xi_i / \phi \xi_i.
$$

(3)

In Eq. (3), $\text{cov}(A, B)$ denotes the covariance of $A$ and $B$, the angle brackets denote expectation value, and we have decomposed the measured values as $\xi_i = \xi_i + \xi_i'$, $\delta_t = \delta_t + \delta_t'$, and $(\delta c)_t = (dc)_t + (dc)_t'$, where primes denote measurement errors and unaccented variables denote true values. Note that the expectation value here is not an average over the data points, which would involve variations in the true $\delta$ and $c$. It is rather the expectation value for a given data point, i.e. what the value for the data point would be if it were an average of many measurements with the true $\delta$ and $c$ held constant.

It follows immediately from the above decompositions that:
\[(\delta c) \delta c' = \delta' \cdot \delta + \delta' \cdot \delta',\]  

(4)

and so:

\[
(\delta c') \delta c = (\delta' \cdot \delta + \delta' \cdot \delta') = \delta' \cdot \delta + \delta' \cdot \delta' \]

\[
= \delta' \cdot \delta + \delta' \cdot \delta' = \delta' \cdot \delta + \delta' \cdot \delta'
\]

(5)

where the second line follows because none of the variables is correlated with another. Thus the correlation coefficient is given by:

\[r = \frac{\delta' \cdot \delta}{\theta_{\delta_i}}\]  

(6)

So for the Keeling plot, we have \(x_i = 1/c_i\) with uncertainty \(\theta_i\), \(y_i = \delta_i\) with uncertainty \(\eta_i\), and \(r = 0\). For the Miller/Tans plot, we have \(x_i = c_i\) with uncertainty \(\epsilon_i\), \(y_i = \delta_i\) with uncertainty \(\phi_i\), and \(r = \delta_i / \phi_i\).

The approximations in Eqs. (1), (2), and (5) are usually excellent. Despite the precision afforded our Monte Carlo simulations by using \(2.5 \times 10^7\) data points, we detect bias due to these approximations only under extreme circumstances (see Section 4.1).

On a modest 2009-model notebook computer, using the Igor Pro programming language (WaveMetrics, Inc.), 5000 five-iteration York fits to a 5000-point mixing line took 215 seconds (compare OLS at 14 seconds and fitexy at 1410 seconds), while 100,000 such fits to a 20-point mixing line took just 25 seconds (compare OLS at 66 seconds and fitexy at 150 seconds). For small numbers of fits, all of these methods are effectively instantaneous. It is perhaps ironic that for most real-world Keeling or Miller/Tans plots, involving fewer than 100 points, solving the York equations is actually faster than OLS.

### 3.2 Monte Carlo simulations

We tested the York, OLS, ODR, GMR, and fitexy methods using simulated measurements of an isotopic mixing line typical of CO\(_2\) respiration in a forest. In this way, the true line is known and the fit bias can be assessed. Our true mixing line had a source isotopic signature of exactly -25‰, a background \(c\) of 380 ppm, and a background \(\delta\) of -9‰. (A Keeling plot of this line has a slope of 6080 and a \(y\)-intercept of -25.) We simulated measurements of this line for a variety of mixing ratio ranges and measurement precisions. For each set of conditions, we generated 5000 'measured' lines, each comprising 5000 points spread evenly over the mixing ratio range \(\Delta c\), with normally distributed, uncorrelated errors added to \(c\) (with standard deviation \(\epsilon\)) and to \(\delta\) (with standard deviation \(\eta\)). The same values of \(\epsilon\) and \(\eta\) were used for all points. We then expressed each line as a Keeling plot and as a Miller/Tans plot, and fit each plot by each of the methods.

Real-world mixing line plots are not likely to contain 5000 points each, but using a large number of points per plot can be important when precisely quantifying fit bias in an ensemble of lines. A demonstration of this point is provided in Appendix B.
We performed tests for $\Delta c = 1, 5, 10, \text{ and } 50$ ppm, and for various combinations of $\varepsilon$ (ppm) and $\eta$ (‰): $\varepsilon = 0.01, \eta = 0.01$ (slightly better than the best existing laser spectrometer, Wehr et al. 2013); $\varepsilon = 0.2, \eta = 0.3$ (for a popular commercial laser spectrometer); $\varepsilon = 0.05, \eta = 0.05$ (for some pairings of an infrared gas analyzer (IRGA) and an isotope ratio mass spectrometer (IRMS)); and $\varepsilon = 0.15, \eta = 0.01$ (for other IRGA/IRMS pairings). Because Kayler et al. (2010) were concerned with biases at very large $\Delta c$ and $\varepsilon$ in studies of leaf respiration and the like, we also conducted simulations for $\Delta c \geq 100$ ppm with $\eta = 0.2$ ‰ as in Kayler et al. (2010) and with $\varepsilon = 1, 5, \text{ and } 20$ ppm.

### 3.3. Forest air measurements

In addition to our Monte Carlo simulations, we analyzed 429 Keeling plots composed of real measurements, specifically nighttime measurements of the mixing ratio and $^{13}$C composition of CO$_2$ in the air within and above a forest canopy. The intercept of such a Keeling plot should give the isotopic composition of nighttime respiration. The measurements were made at each of 6 or 7 heights on a 29 m tower every 40 or 45 minutes from May through October of 2011, 2012, and 2013, as described elsewhere (Wehr et al., 2013; Wehr & Saleska, 2015). Each Keeling plot was made from one night’s data and included about 50 points. The measurement precisions were $\varepsilon = 0.05$ ppm and $\eta = 0.02$ ‰. We also did an alternate analysis in which additional random noise of 0.2 ppm and $\eta = 0.3$ ‰ (corresponding to more common spectroscopic instrumentation) was added to the measurements.

### 4 Results and discussion

#### 4.1 Comparison of fit biases using Monte Carlo simulations

Isotopic source signatures retrieved from our simulated Keeling plots for $\Delta c \leq 50$ ppm are reported in Table 1. The main cells contain three numbers: the upper is the York result (in bold), while the middle is the OLS result, and the lower is the GMR result. We have not tabulated the fitexy or ODR results; the fitexy results are discussed below, and it is enough to say that the ODR results were no better than the GMR results (ODR being a seriously flawed precursor to GMR, as explained in Section 2.1). The units of ‰ here (and throughout this paper) are simply the units of $\delta$, and not an indication of relative error in the results. Numbers in parentheses represent the standard error in the last digit, calculated from the distribution of retrieved values rather than by York’s equations, although the two results did agree closely (see Section 4.2). We have omitted our Miller/Tans results from Table 1 because there were no significant differences between the Miller/Tans and Keeling results for the York and OLS methods, and because the GMR results were very poor for both plot types.

Indeed, as reported by Zobitz et al. (2006), GMR produces highly biased fit lines (that is, the retrieved source signature falls much more than 3 standard errors from -25 ‰). For CO$_2$ ranges less than 50 ppm, the GMR bias is non-negligible in practical terms ($> 0.1$ ‰) unless the measurement uncertainty in $\delta$ is extraordinarily low ($\leq 0.01$ ‰). OLS does better than GMR in most of the tested circumstances because the relative error in the $\varepsilon$ measurement is usually much less than the relative error in the $\delta$ measurement, but OLS still involves important levels of bias when $\varepsilon$ is large or $\Delta c$ is small.
The York equations, on the other hand, produce unbiased Keeling and Miller/Tans fit lines for all conditions in the table. Because the emergence of high-frequency isotopic measurements is starting to raise the issue, we show in detail how some OLS- and York-retrieved isotopic source signatures compare at the lowest Δc in Figure 2, where the error bars represent ±2σ, i.e. twice the standard error in the mean of 5000 fits.

Isotopic source signatures retrieved from our simulated Keeling and Miller/Tans plots for Δc ≥ 100 ppm are reported in Table 2. Again, the York method is by far the best, although the York fit lines do exhibit small but detectable biases for some sets of conditions. The York Miller/Tans fits are biased by at most -0.020 %, while bias in the York Keeling fits is worse, reaching -0.204 % when ε = 20 ppm (an exceptionally high value) and Δc = 100 ppm. This bias is still an order of magnitude less than the bias from any of the other methods under those conditions. The York Miller/Tans bias is due to the approximations made in Eqns. (1) and (5). We have confirmed this by comparing simulations with and without the approximations (a luxury not available with real data). The Keeling bias is due partly to the approximation in Eq. (2), but mostly to the non-normal error distribution in 1/ε (see Section 3.1): a distribution that is asymmetric, with a non-zero mean. In Table 1, where ε (maximum 0.15 ppm) is always a small fraction of c (380 ppm), the skew of the error in 1/ε is small and its effect on the fit negligible; however, in Table 2, where ε/c can be as large as 5 %, the skew becomes relatively large (the uncertainty on one side of a point is about 10 % larger than on the other side) and its effect on the fit becomes detectable in our simulations. The bias induced in the fit by the non-normal error distribution should increase as ε increases and as Δc decreases, which are the trends we observe. The preceding explanation is confirmed by the fact that when we alter our simulations by adding normally distributed measurement error directly to 1/ε rather than to c (and giving the correct information to the York fitting algorithm), we find that the York Keeling fits are completely unbiased (results not tabulated). Luckily, the York fit biases we report in Table 2 are very small considering the measurement uncertainties necessary to induce them, and are unlikely to be the limiting source of error in any experiment.

We also tested fitexy using our Monte Carlo simulations. As expected, the fitexy results were always identical to the York results when fitting to Keeling plots but were in error when fitting to Miller/Tans plots, because the latter plots involved correlation between the x and y errors. For example, with the fairly large measurement uncertainties, ε = 0.2 ppm and η = 0.3 %, the fitexy Miller/Tans slopes were biased by -4.259 % for Δc = 1 ppm and -0.027 % for Δc = 10 ppm.

### 4.2 Comparison of fit biases using real measurements

The intercepts from OLS, GMR, and York fits to our measured Keeling plots are compared in Figure 3, as a function of the CO₂ range. Given that our Monte Carlo simulations show the York fit to be unbiased, we can use the difference between the OLS (or GMR) and York intercepts as a proxy for the bias in OLS (or GMR). In agreement with our Monte Carlo results, Figure 3 shows that for the original measurement uncertainties of 0.05 ppm and 0.02 %, GMR is negligibly biased (that is, by less than 0.1 %) only for CO₂ ranges above about 25 ppm, while OLS is negligibly biased for all CO₂ ranges. Also in agreement with our Monte Carlo results, the figure shows that if the measurement uncertainties are increased to the more common values of 0.2 ppm and 0.3 %, then GMR is never negligibly biased, while OLS is negligibly biased only for CO₂ ranges above 10 ppm. Note that the scatter in Figure 3 is due to the fact that unlike our simulated data points, our real measured data points were not evenly distributed throughout the CO₂ range, in some of the measured Keeling plots, almost all of the points were clustered in a small portion of the range, leading to a higher bias.
4.3 Estimating the fit errors and the goodness of fit

York et al. (2004) provide compact equations not only for the slope and intercept, but for their standard errors as well. They further show that these error estimates are algebraically identical to those of the more general error formulation of MLE. Note, however, that while the York equations for the slope and intercept are exact (if the measurement uncertainties are normally distributed), the York/MLE estimates of the errors in the slope and intercept are approximations (Titterington and Halliday, 1973). In Table 3, we compare York’s error estimates against the standard deviations of the Keeling plot intercepts retrieved from our Monte Carlo simulations. York’s error estimates agree extremely well with the Monte Carlo results except when the measurement error variances are so large as to approach the total variances in x and y (i.e. when $\varepsilon/\Delta c$ and $\eta/\Delta \delta$ approach 1).

Under those conditions, the agreement is nonetheless within 33%.

The errors estimated by fitexy (not tabulated) were slightly higher than those estimated by the York equations, tending to be farther from the Monte Carlo results for small $\varepsilon/\Delta c$ (when the line is well-constrained) and closer to the Monte Carlo results for large $\varepsilon/\Delta c$ (likely by coincidence). The York and fitexy estimated errors were of the same order of magnitude, however, and their disagreement may relate to a subtle point raised in York et al. (2004) concerning whether the errors are estimated using the original data points or what York calls the ‘adjusted’ data points, which are the fit method’s reconstruction of the true, error-free data points.

The standard error is a measure of precision; it does not speak to how well the straight-line model represents the data—a concept known as goodness of fit. York et al. (2004) note that the goodness of fit of the York solution is estimated by the quantity $S/(R-2)$, where $R$ is the number of points in the fit and $S$ is given by:

$$S = \sum W_i (y_i - b x_i - a)^2. \tag{7}$$

This goodness of fit metric is a weighted reduced chi-squared statistic, which we denote here by $\chi^2_0$, i.e. $\chi^2_0 = S/(R-2)$. $\chi^2_0$ is essentially comparing the deviations of the points from the fit line to the assigned measurement error standard deviations. If the variables x and y are in fact related by a straight line, and if the assigned measurement errors are correct (and normally distributed), then $\chi^2_0$ will equal 1. A value of $\chi^2_0$ significantly different from 1 indicates the failure of one or both of those assumptions, where we suggest that significance be defined relative to the standard error in $\chi^2_0$, which depends only on the number of points per fit, R, and is given by:

$$\sigma_x = \sqrt{2/(R-2)}. \tag{8}$$

We have tested this equation by Monte Carlo simulation and found it to be correct to within the simulation precision of roughly 1 part in 200.

The right-hand column of Table 3 gives the mean value of $\chi^2_0$ for each 5000-line ensemble described in the table. With $R = 5000$, our $\sigma_x$ equals 0.02, giving a standard error in our mean $\chi^2_0$ of 3×10^{-4}. Even at this level of precision, $\chi^2_0$ indicates that our fits are almost always good, as expected given the nature of our simulations. However, under those conditions for which the non-normal distribution of 1/$\varepsilon$ is a source of detectible bias, i.e. when $\varepsilon = 20$ ppm, we see the expected deviation in $\chi^2_0$, which
reaches a low of 0.986. In practice, without many fit lines of many points each, such a small drop in $\Delta X\delta$ would be undetectable, as would the associated slope and intercept biases.

### 4.4 Another application of York’s solution: comparing two instruments

When comparing measurements of the same quantity by two different instruments, it is common to plot the values obtained by one instrument against those obtained by the other, so that the relative bias between the instruments can be determined from a straight line fit to the plot. Monte Carlo simulations similar to those used for our isotopic mixing line example confirm that OLS and GMR may incorrectly estimate that bias. For example, if an old, unbiased instrument is being replaced by a new, also-unbiased instrument whose precision is 5 times better, and if the two instruments are compared over a trial period in which the measured quantity varies over a range that is 20 times greater than the precision of the old instrument, then OLS (GMR) will incorrectly indicate that the new instrument is biased low by 4% (2%) of the reading. The York equations will correctly indicate no relative bias.

### 5 Conclusion

We have shown that the general least-squares solution for the best-fit straight line, published by Derek York in 1969, is the least biased estimator of the isotopic signature of a trace gas source from a Keeling or a Miller/Tans plot, regardless of the measurement conditions. In contrast, the popular regression methods considered in the literature are unbiased only under particular, often unrealistic conditions. The isotopic mixing line illustrates the virtue and convenience of York’s solution, which is applicable to line fitting problems in many scientific disciplines. We have provided a short, fast pseudo-code algorithm for computing York’s solution, and derived simple equations for the required measurement error and correlation parameters in the case of a Keeling or Miller/Tans plot. Being both accurate and convenient, York’s solution supersedes all other least-squares straight-line fit methods.

### Appendix A: Computer pseudo-code algorithm

Here we provide an algorithm in pseudo-code for computing the slope and intercept of the best-fit straight line according to Eqs. (13) of York et al. (2004). The data consist of $R$ data points indexed by the subscript $i$. The inputs are: the abscissa $X_i = x_i$; the ordinates $Y_i = y_i$; their measurement error standard deviations $\sigma_{X_i}$ and $\sigma_{Y_i}$; the correlation coefficient between those errors, $r_i$; and an initial guess slope $b_0$. The outputs are: the intercept $a$ and slope $b$, their standard errors $\sigma_a$ and $\sigma_b$, the goodness of fit $\chi^2$, and its standard error $\sigma_{\chi^2}$. The algorithm iterates until the slope converges to within the tolerance $T$. The first block computes the slope and intercept, after which the second block computes the fit errors and the goodness of fit.

```plaintext
b = b0
While $|b_{old} - b| > T$ do
    Begin loop
    $b_{old} = b$
    $X = 0 \& \bar{Y} = 0 \& W_{sum} = 0$
    For $i = 1, R$ step 1 do
```
Begin loop
\[ \omega_{x_j} = 1 / \sigma_{x_j}^2 \quad \text{and} \quad \omega_{\rho_j} = 1 / \sigma_{\rho_j}^2 \]
\[ a_i = \sqrt{\omega_{x_i} \omega_{\rho_i}} \]
\[ W_i = a_i^2 / \left( b^2 \omega_{x_j} \right) + \omega_{x_i} - 2b \rho a_i \]
\[ \bar{X} = \bar{X} + W_i X_j \quad \text{and} \quad \bar{F} = \bar{F} + W_i \rho, \]
\[ \bar{W}_{\text{sum}} = \bar{W}_{\text{sum}} + W_i \]
End loop
\[ \bar{X} = \bar{X} / W_{\text{sum}} \quad \text{and} \quad \bar{F} = \bar{F} / W_{\text{sum}} \]
\[ Q_1 = 0 \quad \text{and} \quad Q_2 = 0 \]
For \( i = 1, R \) step 1 do
Begin loop
\[ U_i = X_i - \bar{X} \quad \text{and} \quad V_i = \rho_i - \bar{F} \]
\[ \beta_i = W_i \left[ (U_i / \omega_{x_j}) + (b \rho_i / \omega_{x_j}) - 2b U_i / \omega_{x_j} \right] \]
\[ Q_i = Q_1 + W_i \beta_i V_i \quad \text{and} \quad Q_2 = Q_2 + W_i \beta_i U_i \]
End loop
\[ b = Q_1 / Q_2 \]
\[ b_{\text{diff}} = |b - b_{\text{old}}| \]
End loop
\[ a = \bar{F} - b \bar{X} \]
\[ \bar{x} = 0 \]
For \( i = 1, R \) step 1 do
Begin loop
\[ x_i = \bar{x} + \beta_i \]
\[ \bar{x} = \bar{x} + W_i x_i \]
End loop
\[ \bar{x} = \bar{x} / W_{\text{sum}} \]
\[ \sigma_b = 0 \quad \text{and} \quad x_b^2 = 0 \]
For \( i = 1, R \) step 1 do
Begin loop
\[ u_i = x_i - \bar{x} \]
\[ \sigma_b = \sigma_b + W_i u_i^2 \]
\[ x_b^2 = x_b^2 + W_i (V_i - b X_j - a)^2 \]
End loop
\[ \sigma_b = \sqrt{\sigma_b^2} \]
\[ \sigma_a = \sqrt{\sigma_a^2 + 1 / W_{\text{sum}}} \]
\[ x_b^2 = x_b^2 / (R - 2) \]
\[ \sigma_x = \sqrt{2 / (R - 2)} \]
Appendix B: Biased estimators and the sample size effect

Why simulate 5000 points per plot when real plots are more likely to contain 20 or 50 points? Consider the simple picture in Figure 4. Here a true line with slope 1 and y-intercept 0 passes through the points (2,2) and (6,6). Measurements at those two points contain errors in \( y \) only, and the errors are supposed to be either +1 or -1. The two measured lines shown have errors that are symmetric in the sense that they are equal in magnitude and opposite in sign, and yet these measured lines are not symmetric about the true line in the sense of being reflections of one another in the true line. Such symmetry would arise only if there were equal error in both \( x \) and \( y \). Nonetheless, the measured slopes (1/2 and 3/2) and y-intercepts (2 and -2) are indeed symmetric about the true values. Now imagine reflecting this picture about \( y = x \) (or rotate the page 90°), so that the line is measured with errors of ±1 in \( x \) only. In this case, the measured slopes (2/3 and 2) and y-intercepts (4/3 and -4) are highly asymmetric about the true values although the fit lines are just as good as before. Instead, the \( x \)-intercepts and inverse slopes have become the symmetric parameters. All this follows from the asymmetries inherent to the definitions of the slope and intercept.

Translating the simple picture of Figure 4 into practical reality, one finds that when there is normally distributed measurement error only in \( y \), straight line fits can be characterized by their slopes and y-intercepts, which are normally distributed parameters that can be averaged over an ensemble of fits to give an unbiased estimate of the slope and y-intercept of the mean fit line (by which we mean the line that would be obtained by combining all of the measured data from the whole ensemble on one plot and doing a single fit to that). However, when error is present also (or only) in \( x \), the slopes and y-intercepts of an ensemble of fit lines are not normally distributed about the mean-line values. Instead the distributions are skewed, with means and modes that differ from one another and from the mean-line values. In the general case, with uncertainty in both \( x \) and \( y \) that might vary from point to point, there are no standard fit line parameters that can be averaged to give the corresponding parameters of the mean fit line.

Fortunately, as the CO\(_2\) range or the number of points per fit line becomes large, the deviations of the fit lines from the true line become small and the skew in the parameter distributions becomes insignificant. We exploited this fact in our Monte Carlo simulations by using 5000 points per line, which enabled us to simply average our fit slopes and intercepts without introducing significant bias even for \( \Delta c = 1 \) ppm. Real-world Keeling plots often contain only 10 or 20 points, but usually span more than 10 ppm, and so the skew remains small. Moreover, real-world Keeling plots are not found in ensembles of 5,000, and so the statistical error in the mean slope and intercept would swamp any skew-related bias even for \( \Delta c = 1 \) ppm.

One might be worried about increasing the number of points per plot because it has been reported that fit bias appears to increase with the number of points per plot (Kayler et al., 2010). Actually, increasing the number of points per plot merely clarifies the bias associated with a poor fitting method. Consider Figure 5, where we plot distributions of y-intercepts retrieved from Keeling plots with \( \Delta c = 1 \), \( \varepsilon = 0.15 \), and \( \eta = 0.01 \). These conditions are extreme in that \( \varepsilon / \Delta c \approx \eta / \Delta \delta \approx 0.2 \) and so the measurement error spread in each dimension is roughly equal to the extent of the true line. Effectively similar conditions (\( \Delta c = 40 \), \( \varepsilon = 5 \) ) were used to produce Table 3 of Kayler et al. (2010). In our figure, the thick curve is the distribution with just 2 points per line, in which case there is no distinction between fit methods because “fitting” consists merely of drawing a line exactly between the two measured points. As the number of points per line increases, they form a distribution in Cartesian space that each fit method interprets differently (according to its assumptions about the measurement error distributions and correlation), so that the intercept distributions obtained from different fit methods diverge. The thin solid curves in Figure 5 show the distributions retrieved from the York equations as the number of points per plot increases to 5 and then 20. The dashed curves show the
corresponding distributions retrieved from OLS. By 20 points, the distributions are nearly Gaussian, and the York distribution is nearly centered on the true mean while the OLS distribution is nearly centered on the value corresponding to its true fit bias under these uncertainty conditions. Note that the skew in Figure 5 is not due to the subtly skewed noise distribution in the abscissa: the figure is not visibly changed when normally distributed noise is added directly to 1/c rather than to c.

5 Author contributions

R. Wehr designed and performed the study, and prepared the manuscript with contributions from S. R. Saleska.

Competing interests

The authors declare that they have no conflict of interest.

Acknowledgments

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References


Pearson, K.: On lines and planes of closest fit to systems of points in space. Philos. Mag., 2(6), 559–572, 1901.


Figures, Tables, and Captions

![Figure 1: An example Keeling plot from the set described in Section 3.3, showing 55 measurements made on the night of May 25, 2011, spanning a CO₂ range Δc = 5 ppm, with measurement error standard deviations of $\sigma = 0.05$ ppm and $\eta = 0.02\%$.](image-url)
Figure 2: Mean isotopic signatures retrieved from ensembles of 5000 simulated Keeling plots (each containing 5000 points) using the York and OLS methods, for CO₂ ranges from 1 to 10 ppm and a true isotopic signature of -25 ‰. The standard deviations of the random noise added to the simulated measurements of ε and δ are inset. Error bars show two standard errors. Note the differing scales for the ordinate.

Figure 3: Difference between the fit intercept obtained by OLS or GMR and that obtained by York’s equations, for 429 real measured Keeling plots with measurement uncertainties of 0.05 ppm and 0.02 ‰ and with CO₂ mixing ratio ranges between 2 and 335 ppm. Also shown is the difference obtained for Keeling plots in which random computer-generated noise of 0.2 ppm and 0.3 ‰ was added to the original data. The grey region represents biases that are negligible for most practical purposes (< 0.1 ‰).
Figure 4: Two lines, each defined by measurements of the true points (2,2) and (6,6) that are in error by +1 or -1 in the ordinate only.

Figure 5: Distributions of the $y$-intercepts retrieved from Keeling plots with a true intercept of -25‰ (indicated by the vertical line). The black curve was obtained from $10^6$ two-point plots. The red curves were obtained using the York equations on $10^5$ five-point or twenty-point plots. The blue curves were obtained using OLS on $10^5$ five-point or twenty-point plots. The heights of the curves have been adjusted independently for presentation and convey no meaning.
Table 1. Bias in retrieved isotopic source signatures from Keeling line fits for Δc ≤ 50 ppm, expressed as differences from the true value of -25, in units of ‰. Each cell contains, from top to bottom: the York, the OLS, and the GMR result. 1-σ uncertainties in the final digit(s) are given in parentheses, based on the standard deviation of the ensemble of 5000 fits.

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<th>η (%o)</th>
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<th>10</th>
<th>50</th>
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Table 2. Bias in retrieved isotopic source signatures from Keeling and Miller/Tans line fits for $\Delta c \geq 100$ ppm and $\eta = 0.2 \, \%$, expressed as differences from the true value of -25, in units of $\%$. Each cell contains, from top to bottom: the York, the OLS, and the GMR result. 1-$\sigma$ uncertainties in the final digit are given in parentheses, based on the standard deviation of the ensemble of 5000 fits.

<table>
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Comment [6]: Table has been rearranged, dropping out the 5000 ppm column.
Table 3. Monte Carlo (MC) and York estimates of the error in the isotopic source signature retrieved from an individual Keeling plot, in units of ‰. The mean goodness of fit $\chi^2_\nu$ over each 5000-fit ensemble is also shown.

<table>
<thead>
<tr>
<th>$\Delta c$</th>
<th>$\epsilon$ (ppm)</th>
<th>$\eta$ (%)</th>
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<th>York</th>
<th>$\chi^2_\nu$</th>
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