This report describes mathematical relationships between step and cumulative process mass intensities (PMIs) for synthesis plans, and analogous parameters applied to E-factors. It is shown that both step E-factors and step PMIs are not additive for synthesis plans. It is also shown that a recursive calculation of cumulative PMIs from step PMIs is a rapid method of determining overall PMIs for synthesis plans, though cumulative PMIs are not sufficiently informative as step PMIs or step E-factors to identify bottlenecks in synthesis plans. Illustrations on the use of these metrics to track the material efficiency of published synthesis plans for the pharmaceutical, apixaban, are given as a template example. Advantages and disadvantages of each metric are discussed. A general algorithm to select the most promising candidate synthesis plans considered at the design stage for a given molecular target that most likely satisfy “green” material efficiency criteria is also presented.
Relationships Between Step and Cumulative PMI and E-factors: Implications on Estimating Material Efficiency With Respect to Charting Synthesis Optimization Strategies

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Abstract:
This report describes mathematical relationships between step and cumulative process mass intensities (PMIs) for synthesis plans, and analogous parameters applied to E-factors. It is shown that both step E-factors and step PMIs are not additive for synthesis plans. It is also shown that a recursive calculation of cumulative PMIs from step PMIs is a rapid method of determining overall PMIs for synthesis plans, though cumulative PMIs are not sufficiently informative as step PMIs or step E-factors to identify bottlenecks in synthesis plans. Illustrations on the use of these metrics to track the material efficiency of published synthesis plans for the pharmaceutical, apixaban, are given as a template example. Advantages and disadvantages of each metric are discussed. A general algorithm to select the most promising candidate synthesis plans considered at the design stage for a given molecular target that most likely satisfy “green” material efficiency criteria is also presented.

Keywords: green chemistry education, green chemistry engineering, sustainability metrics

Abbreviations:
AE = atom economy
BP = by-products
CAT = catalyst
cE = cumulative E-factor
cPMI = cumulative process mass intensity
\[ E = \text{E-factor} \]
\[ m = \text{mass} \]
\[ \text{MW} = \text{molecular weight} \]
\[ N = \text{number of reaction steps} \]
\[ \text{PMI} = \text{process mass intensity} \]
\[ \text{PU} = \text{purification} \]
\[ \text{RS} = \text{reaction solvents} \]
\[ \text{SP} = \text{side products} \]
\[ \text{WU} = \text{work-up} \]
\[ \text{XS} = \text{excess} \]

Greek letters:
\[ \varepsilon = \text{reaction yield} \]
\[ \phi = \text{mass of input materials} \]

**Introduction:**

The use of green chemistry metrics is now a well-established tool used by synthetic organic and process chemists to estimate the material efficiency, environmental impact, and safety-hazard impact of their synthesis plans and chemical processes to a desired target molecule \([1,2]\). More importantly, it is a powerful tool in making decisions about which candidate synthesis strategies to choose and ultimately to actualize in the pursuit of “green” syntheses at the early design and planning stages of a synthesis campaign. Even more importantly, it is imperative that whatever algorithms are used to carry out the estimation of those metrics are robust so that good decisions are made with respect to achieving true synthesis optimization. In this way, chemists will be confident in implementing reliable, reproducible, and intelligible metrics as a routine tool at their disposal in their practice of green chemistry principles in their everyday work.

In this report we wish to present general relationships between step and cumulative process mass intensity (PMI) and general relationships between step and cumulative E-factors for linear and convergent synthesis plans. Both sets of metrics have been advanced by the pharmaceutical industry as highly relevant to the determination of material efficiency both for “after-the-fact” published plans in the literature and for exploring new synthesis strategies using classical and novel methods at the planning
stages before any experiments are carried out. Specifically, we develop facile methods of calculating overall PMI, cumulative PMI, overall E-factor, and cumulative E-factor for synthesis plans. The structure of this report is as follows. We begin by showing the derivation of each method and supply mathematical evidence supporting the connections between the relevant metrics. Next, we illustrate the computation of those metrics for published synthesis plans to the blood anti-coagulant pharmaceutical apixaban 1 [3-8] since this was chosen as a showcase example by the pharmaceutical industry for achieving a “green” synthesis [9]. Finally, we show how these metrics may be used for ranking the published plans with respect to material efficiency.

**E-factor and PMI**

For any individual chemical reaction, the mass of waste generated may be generally considered to originate from seven sources: by-products (BP) arising as a mechanistic consequence in producing the target product, side products (SP) arising as a consequence of competing reactions not leading to the intended target product, unreacted excess reagents (XS), catalysts and ligands (CAT), reaction solvents (RS), work-up materials (WU), and purification materials (PU). If $m_p$ is the mass of the desired target product of the reaction, then the total mass of waste and the E-factor are defined according to equations (1) and (2) shown below.

$$m_{\text{waste}} = m_{BP} + m_{SP} + m_{XS} + m_{CAT} + m_{RS} + m_{WU} + m_{PU}$$  \hspace{1cm} (1)

$$E = \frac{m_{\text{waste}}}{m_p} = \frac{m_{BP} + m_{SP} + m_{XS} + m_{CAT} + m_{RS} + m_{WU} + m_{PU}}{m_p}$$

$$= \frac{m_{BP}}{m_p} + \frac{m_{SP}}{m_p} + \frac{m_{XS}}{m_p} + \frac{m_{CAT}}{m_p} + \frac{m_{RS}}{m_p} + \frac{m_{WU}}{m_p} + \frac{m_{PU}}{m_p}$$  \hspace{1cm} (2)

$$= E_{BP} + E_{SP} + E_{XS} + E_{CAT} + E_{RS} + E_{WU} + E_{PU}$$

We observe from equation (2) that the E-factor for the reaction can be decomposed into its contributing E-factors based on the individual sources of the waste. Hence, the overall E-factor for a single chemical reaction is the sum of the contributing E-factors corresponding to each waste source. Now, if we follow a parallel manipulation using the
corresponding PMI parameters, then we need to use the general connecting relationship \( PMI = E + 1 \) \[10\]. For an individual reaction where \( m_{\text{input}} \) represents the mass of all input materials used, the expression for PMI can be written as shown in equation (3).

\[
PMI = \frac{m_{\text{input}}}{m_p} = \frac{m_{\text{react}} + m_p}{m_p} = \frac{m_{\text{waste}}}{m_p} + 1 = E + 1
\]  

(3)

Hence, the last line in equation (2) can be transformed as shown in equation (4).

\[
PMI - 1 = (PMI_{BP} - 1) + (PMI_{SP} - 1) + (PMI_{XS} - 1) + (PMI_{CAT} - 1) \\
+ (PMI_{RS} - 1) + (PMI_{WU} - 1) + (PMI_{PU} - 1)
\]

\[
PMI = PMI_{BP} + PMI_{SP} + PMI_{XS} + PMI_{CAT} + PMI_{RS} + PMI_{WU} + PMI_{PU} - 6
\]  

(4)

Equation (4) indicates that the overall PMI for a chemical reaction is not the sum of the contributing PMIs as was the case for E-factors. Rather, it is the sum of the contributing PMIs minus one less than the number of waste contributing factors. Based on these findings for a single reaction, one might think that the same kind of parallel relationship also holds for sequences of reactions in a synthesis plan; namely, that the overall E-factor is the sum of the step E-factors and that the overall PMI is not the sum of the step PMIs. In fact, for a linear synthesis plan neither the overall E-factor nor the overall PMI is the sum of the corresponding step parameters. We demonstrate what the connecting relationships are between overall and step parameters for each metric with the following simple logic. For a linear sequence of \( N \) steps involving the production of intermediate products \( P_1, P_2, \ldots, \) and \( P_N \) in steps 1, 2, \ldots, \( N \) where all of the preceding intermediate in any given step is committed as a reagent in the next step, we can write the step PMI for step \( j \) as shown in equation (5).

\[
(PMI)_j = \frac{\phi_j}{m_{p_j}},
\]  

(5)

where \( \phi_j \) is the mass of input materials used in step \( j \) and \( m_{p_j} \) is the mass of intermediate product collected in step \( j \). For the entire synthesis plan covering all \( N \) steps, the overall PMI is given by equation (6) where we have substituted the step PMI expression given in equation (5).
From equation (6) we observe immediately that the overall PMI is generally not the sum of the step PMIs. However, the expression can collapse to that sum under the very special condition when all the masses of the intermediate products are each identical to the final product of the linear synthesis, i.e., $m_{P_j} = m_{P_N}$ for $j = 1, 2, 3, \ldots, N$. Such a scenario implies a linear sequence comprising consecutive rearrangement reactions where the molecular weight of each intermediate product remains the same and each reaction yield is 100% with no loss of mass arising from by-products or unreacted starting materials along the way. This very rare possibility involving consecutive rearrangement reactions has been documented before [11]. We can transform the expression given in equation (6) to find an analogous expression given in equation (7) that relates the overall E-factor for a linear sequence to the step E-factors using the substitution $PMI = E + 1$.

$$E_{i \rightarrow N} + 1 = \frac{1}{m_{P_N}} \left[ m_{P_N} (E_N + 1) + \sum_{j=1}^{N-1} m_{P_j} \left( (E_j + 1) - 1 \right) \right]$$

$$= \frac{1}{m_{P_N}} \left[ m_{P_N} (E_N + 1) + \sum_{j=1}^{N-1} m_{P_j} E_j \right]$$

$$E_{i \rightarrow N} = E_N + \frac{1}{m_{P_N}} \sum_{j=1}^{N-1} m_{P_j} E_j$$

Again, we observe that the overall E-factor is generally not the sum of the step E-factors, except for the very special case mentioned above. This mathematical evidence therefore shows that step E-factors for a linear plan are not additive. Close examination of equation (7) shows that each of the E-factor parameters is defined with respect to
different product masses. Now, if we define the overall mass of waste produced in a linear synthesis plan as the difference between the sum of masses of all input materials used in all reaction steps and the sum of masses of all products produced along the way as shown in equation (8)

\[ m_{\text{overall \ waste}} = \sum_{j=1}^{N} \phi_j - \sum_{j=1}^{N} m_{P_j} \]  

(8)

and then if we divide that mass of overall waste by the mass of the final target product we obtain, of course, the overall E-factor for the synthesis plan. Furthermore, if we define a step contributing E-factor, \( E_j^* \), with respect to the mass of final target product representing the contribution of waste produced from each reaction step to the total waste produced, then we can transform the expression given in equation (8) to obtain a relationship that links the overall E-factor to the step contributing E-factors as shown in equation (9).

\[
\frac{m_{\text{overall \ waste}}}{m_{P_N}} = \frac{\phi_1 - m_{P_1}}{m_{P_N}} + \frac{\phi_2 - m_{P_2}}{m_{P_N}} + \ldots + \frac{\phi_N - m_{P_N}}{m_{P_N}} 
\]

\[
E_{\text{overall}} = \frac{1}{m_{P_N}} \left( E_1^* + E_2^* + \ldots + E_N^* \right) 
\]

In this case, we see that, indeed, the overall E-factor for a linear synthesis plan is the sum of the step contributing E-factors. Note that each step E-factor appearing in equation (8) is defined with respect to the mass of intermediate product produced at that step, whereas the step contributing E-factors appearing in equation (9) are all defined with respect to the mass of final product in the sequence.

**Cumulative Versus Step Metrics**

We previously reported expressions for cumulative PMI and cumulative E-factor that depended on step PMI and step E-factors, respectively [12,13]. The recursive relationships for linear sequences are shown in equations (10) and (11).
\[(cPMI)_{1\rightarrow i} = \frac{m_{PMI}}{m_i} \left[ (cPMI)_{1\rightarrow i-1} - 1 \right] + (PMI)_i \]  
\[(10)\]

\[(cE)_{1\rightarrow i} + 1 = \frac{m_{PMI}}{m_i} \left[ (cE)_{1\rightarrow i-1} + 1 - 1 \right] + (E)_i + 1\]

\[(cE)_{1\rightarrow i} = \frac{m_{PMI}}{m_i} (cE)_{1\rightarrow i-1} + (E)_i\]  
\[(11)\]

where \( \frac{m_{PMI}}{m_i} = \frac{MW_{PMI}}{MW_i \epsilon_i} \), \( MW \) refers to molecular weights of intermediates multiplied by stoichiometric coefficients as appropriate from balanced chemical equations, \( \epsilon_i \) is the reaction yield of step \( i \), the \( 1 \rightarrow i \) notation means that the cumulative PMI quantity extends from step 1 to step \( i \), and the counting index goes from \( i = 2, 3, \ldots, \) etc. Note that the connecting relation \( PMI = E + 1 \) was used in obtaining equation (11) from equation (10). The derivation of equation (10) arises as follows. For a three-step plan, from equation (5), the step PMIs for steps 1, 2, and 3 are given by equations (12-14).

\[(PMI)_1 = \frac{\phi_1}{m_i}\]  
\[(12)\]

\[(PMI)_2 = \frac{\phi_2}{m_i}\]  
\[(13)\]

\[(PMI)_3 = \frac{\phi_3}{m_i}\]  
\[(14)\]

The cumulative PMI for steps 1 and 2 is given by equation (15).
\[ (cPMI)_{1\to2} = \frac{\phi_1 + \phi_2 - m_{P_1}}{m_{P_2}} = \frac{\phi_1}{m_{P_1}} + \frac{\phi_2}{m_{P_2}} - m_{P_1} \]
\[ = \frac{\phi_1}{m_{P_1}} \cdot \frac{m_{P_2}}{m_{P_1}} + \phi_2 - m_{P_2} \cdot \frac{m_{P_1}}{m_{P_2}} = m_{P_1} \left[ \frac{\phi_1}{m_{P_1}} - 1 \right] + \frac{\phi_2}{m_{P_2}} \]
\[ = m_{P_1} \left[ (PMI)_1 - 1 \right] + (PMI)_2 \]
\[ = m_{P_1} \left[ (cPMI)_{1\to1} - 1 \right] + (PMI)_2 \]  

where it is understood that the step PMI for step 1 is identical to the cumulative PMI for step 1. Similarly, the cumulative PMI for steps 1, 2, and 3 is given by equation (16).

\[ (cPMI)_{1\to3} = \frac{\phi_1 + \phi_2 + \phi_3 - m_{P_1} - m_{P_2} - m_{P_3}}{m_{P_3}} = \frac{m_{P_1} (cPMI)_{1\to2} + \phi_3 - m_{P_3}}{m_{P_3}} \]
\[ = m_{P_1} \left[ (cPMI)_{1\to2} - 1 \right] + (PMI)_3 \]  

It is readily apparent from the emerging pattern of equations (15) and (16) that equation (10) is verified for general step \( i \). For a linear sequence consisting of \( N \) steps, the cumulative PMI, \( (cPMI)_{1\toN} \) is identical to the overall PMI that would be obtained from equation (6). It is also understood from this analysis that all intermediate products produced along the linear chain are limiting reagents in each reaction, so the molar stoichiometric ratio is always 1. When we are dealing with a convergent synthesis the same kind of analysis is done on the convergent branch from the first step of that branch to the final step of that branch. A corresponding cumulative PMI will therefore be obtained up to the terminal intermediate of the convergent branch. At the convergent step there will be a reaction between an intermediate product from the main branch and the terminal product of the convergent branch along with any other reagents. In a properly designed synthesis the intermediate product from the main branch will be the limiting reagent, but the terminal intermediate product from the convergent branch may be used in excess. Hence, the expression for the cumulative PMI from step 1 up to the convergent step along the main branch will have an extra term included that accounts for the contribution arising from the convergent branch. For example, if we have a
convergent synthesis plan having a convergent branch consisting of 3 steps leading up to intermediate \( P_3 \), and the convergent step occurs at step 5 along the main branch then the cumulative PMI from step 1 to step 5 is given by equation (17).

\[
(cPMI)_{1 \rightarrow 5} = \frac{m_{P_5}}{m_{P_4}} [(cPMI)_{1 \rightarrow 4} - 1] + (PMI)_5 + \frac{m_{P_5}}{m_{P_4}} [(cPMI)_{1 \rightarrow 3} - 1]
\]  

(17)

The first two terms in equation (17) arise from the normal recursion formula applied to the main branch and the last term pertains to the contribution arising from the convergent branch. The expressions for the two mass ratios of intermediate products are shown in equations (18) and (19).

\[
\frac{m_{P_5}}{m_{P_4}} = \frac{MW_{P_5}}{MW_{P_4} \varepsilon_5}
\]  

(18)

\[
\frac{m_{P_5}}{m_{P_4}} = \frac{MW_{P_5}}{MW_{P_4} \varepsilon_5} \times \frac{\text{actual moles}_{P_5}}{\text{stoichiometric moles}_{P_5}}
\]  

(19)

where the mole ratio in equation (19) is a number that has a value greater than 1.

Appendix 1 contains a more complex example of a convergent plan drawn in the form of a synthesis tree diagram [14,15] containing 5 branches including the main branch (See Figure S1). From that diagram it is possible to write out all the cumulative PMI expressions sequentially from the first step all the way to the final step by inspection (see equations S1 to S17).

**Application to Syntheses of Apixaban**

We evaluated the material efficiency of various literature-documented syntheses of the blood anti-coagulant apixaban \( 1 \) as a demonstration of the mathematical relationships presented in this work. Six synthesis plans were considered including two from Bristol-Myers Squibb [3,4], one from an academic lab in China [5], and three from generic drug companies [6-8]. All of them follow the same convergent synthesis strategy as evidenced by the common target bond mapping shown in Figure 1 which highlights
the construction bonds made in the product chemical structure. The mapping for the pyrazolo[3,4-c]pyridin-7-one ring frame can be encoded as \([(2 + 1 + 1 + 1) + (5 + 1)]\).

**Figure 1**

Table 1 summarizes the essential materials efficiency metrics for all six plans which are listed in ascending order of total PMI. These metrics were calculated using our previously reported REACTION and SYNTHESIS spreadsheets [11]. Best metric parameters are highlighted in green and worst ones are shown in red. This ranking shows that the Bristol-Myers Squibb 2006 plan has the best overall mass efficiency profile having the best performances with respect to highest overall yield and overall atom economy, and the lowest values for E-kernel, E-excess, E-solvent, E-workup, E-total, and PMI-total. That plan corresponds to the one reported in a recent paper on cumulative PMI calculations [9]. The Jiang-Ji plan had the lowest E-cat contribution. The separation of E-total contributions assists in pinpointing each plan’s strengths and weaknesses. The main liabilities impacting the last place DrReddy plan are a high reaction solvent consumption and a high work-up auxiliary material consumption. The significant improvement in the Bristol-Myers Squibb 2006 plan over their earlier 2003 plan is mainly due to an 11% gain in overall yield and reductions in E-solvent and E-workup by 59% and 294%, respectively. It is not surprising that most of the potential gains in “greening up” synthesis procedures arise from solvent and other auxiliary material consumption since these account for at least 75% of all the input mass used in chemical reactions. As a consequence pharmaceutical companies have focused their efforts to reduce auxiliary material consumption as a primary strategy to achieve green syntheses of their products. We can drill deeper to find the precise reaction steps in each plan that are responsible for any mass inefficiencies by examining the shapes of the corresponding radial pentagon diagrams and correlating them to which of the following 4 parameters is responsible: reaction yield, step AE, step SF (stoichiometric factor representing excess energy consumption), and MRP (material recovery parameter representing auxiliary material consumption). Figure 2 shows the radial pentagon diagrams for the Bristol-Myers Squibb 2006 apixaban plan. By visual inspection that
plan has the following liabilities: (a) low AE for steps 1 (amidation-\(\alpha,\alpha'\)-dichlorination sequence) and 2b (diazotization-imation sequence) and (b) high SF for steps 2b and 6 (final carboxyamination). Applying the same analysis to the other plans we have the following results. For the Optimus 2017 plan we have the following liabilities: (a) low AE for steps 2 (cyclic amination), 3 (\(\alpha,\alpha'\)-dichlorination), 5 (Heck coupling), and 5b (diazotization-imation sequence), (b) high SF for steps 5, 5b, and 8 (final carboxyamination). For the Jiang-Ji 2013 plan we have the following liabilities: (a) low AE for steps 1 (cyclic amidation-amination sequence), 2 (\(\alpha,\alpha'\)-dichlorination-morpholine substitution sequence), 3 (aromatic nitro group reduction with sodium sulfide), 4 (cyclic amidation-amination sequence), and 4b (diazotization-imation sequence), and (b) high SF for steps 1, 2, and 4b. For the MSN 2015 plan we have the following liabilities: (a) low reaction yields for steps 2 (cyclic amination-\(\alpha,\alpha'\)-dichlorination-chlorine elimination sequence) and 4b (diazotization-imation sequence), (b) low AE for steps 1 (acid chloride formation-amidation sequence), 2, 4 (Heck coupling), and 4b, and (c) high SF for steps 4b, 5 (pyrazole formation), and 6 (carboxyamination). For the Bristol-Myers Squibb 2003 plan we have the following liabilities: (a) low reaction yields for steps 2 (\(\alpha,\alpha'\)-dichlorination-morpholine substitution sequence) and 5 (final carboxyamination), (b) low AE for steps 1 (amidation-cyclic amination sequence), 2, 3 (Heck coupling), and 2b (imation), and (c) high SF for steps 2, 4 (pyrazole formation), and 5. Finally, for the DrReddy 2017 plan we have the following liabilities: (a) low reaction yield for step 3 (pyrazole formation), (b) low AE for steps 1 (\(\alpha,\alpha'\)-dichlorination), 2 (chlorine elimination), 2b (diazotization-imation sequence), 3, 3bb (amidation-cyclic amination sequence), and 4 (Heck coupling), and (c) high SF for steps 2b, 3, and 4.

Figure 2

For comparison we summed the step E-factors for each plan and obtained the following values: 197.81 (Bristol-Myers, 2006); 208.72 (Optimus); 338.30 (Jiang-Ji); 274.18 (MSN); 419.51 (Bristol-Myers, 2003); and 688.55 (DrReddy). We can see that in all cases that the values obtained by summing the step E-factors are noticeably less than the true values of E-total shown in the 11th column of Table 1. The additive values are
found to be between 12 and 45 % lower than the true overall E-factor values. This shows that in this specific case the summing of step E-factors results in a significant underestimation of overall E-factor. However, this result does not always hold true all the time as we demonstrate that there are other cases when the sum of step E-factors may exceed the true overall E-factor. In Appendix 2 we show as examples all possible case scenarios for general two- and three-step synthesis. For longer syntheses it is difficult to make predictions as to the relative outcome of the sum of E-factors versus the true overall E-factor because the number of possible scenario cases increases dramatically making the task impractical. What is certain is that the sum of step E-factors does not equal the true overall E-factor for a given synthesis plan.

Table 1

Following our developed cumulative PMI calculator spreadsheet based on the recursive relation given in equation (10) (see Supporting-Information-EXCEL.zip) we were able to easily obtain the cumulative PMI values at each reaction step for all synthesis plans examined. The input data include the reaction yields, the molecular weights of intermediate products and final product, the mole ratios of terminal intermediate products from the convergent branches, and the step PMIs. The step PMIs were obtained from the REACTION spreadsheets applied to each reaction step using the reported experimental masses of materials used. The results are reported in Tables 2, 3, and 4 corresponding to complete, kernel, and kernel plus excess cumulative PMI values, respectively. The three levels of calculation correspond to how the step PMI values are determined. The complete calculations correspond to all materials used, the kernel calculations correspond to only atom economy and reaction yield, and the kernel plus excess reagent contribution corresponds to atom economy, reaction yield, and stoichiometric factor as shown in equations (20) to (22).

\[
\text{step } PMI_{\text{complete}} = \left[ (Yield)(AE) \left( \frac{1}{SF} \right)(MRP) \right]^{-1} \tag{20}
\]

\[
\text{step } PMI_{\text{kernel}} = \left[ (Yield)(AE) \right]^{-1} \tag{21}
\]
\[
\text{step PMI}_{\text{kernel+excess}} = \left[ \frac{\text{Yield}}{\left( \frac{AE}{SF} \right)} \right]^{-1}
\]
(22)

where \( SF \) is the stoichiometric factor [16], and \( MRP \) is the material recovery parameter accounting for all auxiliary materials used [16].

The highlighted values in yellow shown in Tables 2, 3, and 4 correspond to the overall cumulative PMIs for each plan under the three regimes of calculation. In all cases the results of the final cumulative PMIs agreed with the overall PMIs obtained from the SYNTHESIS spreadsheet analyses. For example, the highlighted values shown in Table 2 agree with those given in the last column of Table 1. Table 5 shows the rankings of all plans based on these overall cumulative PMI values. We can see that the Bristol-Myers Squibb 2006 plan ranks first in all three regimes whereas the ranking order changes for the other plans. This actually is an excellent test for robustness of “green” material efficiency for a synthesis. If a plan consistently ranks first under these three regimes then the probability that it represents the “greenest” plan to a given target is very high. Ranking first under the kernel regime means that the plan likely has the highest atom economy and overall yield; ranking first under the kernel plus excess regime means that it also consumes the least excess reagents; and finally ranking first under the complete regime means that it also consumes the least auxiliary materials in terms of reaction solvents, work-up materials, and purification materials.

Table 2
Table 3
Table 4

A close examination of the rankings of the other plans shown in Table 5 shows some significant changes depending on the calculation regime. For example, the Optimus plan is ranked #2 under the complete regime but drops to #5 under the kernel regime. This is consistent with its auxiliary material consumption being close to that of the #1 plan, and its low ranking overall yield – about half that of the #1 plan. Apart from
the winning plan, the Jiang-Ji and MSN plans show little variation in ranking among the three regimes. It is clear from this example that true overall synthesis optimization is achieved when optimization in all three regimes is orchestrated in the same direction. We believe and emphasize that this criterion is the best and most significant indicator that controls material efficiency optimization of synthesis plans, and therefore the achievement of comparatively “green” synthesis plans. We also point out that in order to drill down to discover where the bottlenecks are in any given plan, it is insufficient to look only at overall E-factor, overall PMI, overall AE, and overall yield values as guides, or for that matter cumulative counterparts of those parameters. Two important guides are the partitioning of E-factors as shown in Table 1, and radial pentagon diagrams for each reaction step in a synthesis as shown in the REACTION spreadsheets for each plan and discussed elsewhere [12]. We determined the precise liabilities of each synthesis plan for apixaban by following this procedure as described in our earlier discussion. We note that the computation of overall PMI via a recursive cumulative PMI calculation is indeed facile. However, the tracking of the incremental increases in PMI does not shed enough light on spotting bottlenecks in a synthesis plan in comparison to tracking individual reaction step PMIs in conjunction with their radial pentagon diagrams which show how the partitioned step metrics come together to result in the corresponding step PMI values.

Table 5

Though it appears that synthesis optimization of apixaban using the existing synthesis strategy has reached a plateau, it does not mean that entirely new strategies cannot be considered that would further drive optimization forward. What has been achieved is comparatively the “greenest” synthesis plan within the constraint of a particular design synthesis strategy as shown by the common target bond mapping shown in Figure 1. We demonstrate that it is possible to use the kernel green metrics of reaction yield and atom economy applied to new synthesis schemes employing new synthesis design strategies to the same target molecule. Once a scheme is conjectured its step and overall atom economies can be determined. Estimates of reaction yield performance at each step in a conjectured scheme may be made via literature searches where minimum
and maximum yield ranges may be obtained. Selecting maximum yield values based on literature evidence coupled with atom economy metrics will yield best-case scenario step PMI and cumulative PMI computations under the kernel regime. These results may be compared with those from the documented literature plans computed under the same regime. In addition, these results may then be judged to see if any of the conjectured schemes have any merit for further investigation pending criteria such as availability and cost of starting materials, environmental impact issues with respect to input materials and by-products produced, safety and hazard issues pertaining to the use of any reagent or the handling of waste reaction by-products, opportunities for recycling and/or reclaiming, and anticipation of unwanted side reactions occurring.

Algorithm for Finding the Optimum Material Efficient Synthesis Plan

Based on our ongoing research spanning 15 years on the study of quantitative parameterization of synthesis and green metrics development, we are able to put forward the following algorithmic protocol for finding the optimum material efficient synthesis route to a desired known target molecule. Two main stages are involved: (i) in stage I a thorough literature search of existing plans is carried out with a view to create a complete reaction network as shown by the generalized example given in Figure 3; and (ii) in stage II, based on the findings of stage I, a brainstorming of ideas is summoned to come up with new synthesis strategies not documented in the literature for the same target molecule. Stage I involves a comprehensive analyses of known synthesis plans from properly documented experimental procedures using step and overall reaction yield, atom economy, E-factor, E-factor partitions, and PMI metrics conducted under the complete and kernel regimes. Stage II involves inventing new synthesis strategies leading to different target bond maps for the same target molecule and assessing their potential for material efficiency “greenness” using the same suite of metrics as Stage I but under the kernel regime.

Figure 3
In the case of the apixaban plans studied in this work, the corresponding reaction network summarizing all intermediates involved in all routes is shown in Figure 4. Figures 5 and 6 show flowcharts describing the logical flow of steps involved in stages I and II. We followed both sets of protocols in presenting our findings on the synthesis of apixaban in this work.

Conclusions:

In this work we have established the mathematical connections between step and cumulative E-factor and PMI metrics with a view to facilitate computation of these quantities for synthesis plans. The statement that the sum of step E-factors for a synthesis plan does not equal the overall E-factor has been confirmed. Similarly, the sum of step PMIs for a synthesis plan does not equal the overall PMI. We also presented a simple computation of cumulative PMI and produced a simple calculator that automates the calculation for both linear and convergent plans. We have demonstrated the utility of a metrics-guided algorithm to both comprehensively assess literature synthesis plans to a given target molecule and to suggest new synthesis strategies for the same product using the synthesis of the pharmaceutical apixaban as an example. We believe that the quantitative parameterization of synthesis performance of individual reactions and synthesis plans (both linear and convergent) by green metrics at the level of material efficiency is now a completely solved problem and does not require further investigation or invention of new metrics or algorithms. The only outstanding problem is deciding how far back one needs to trace intermediates and reagents to a set of progenitor “starting materials” in order to conduct a materials efficiency metrics analysis of a given synthesis plan. One of the tasks of the recently convened IUPAC Metrics for Green Syntheses (MGS-1) Project is to tackle this question [13]. We hope that this work will put chemists on a confident path to optimize their own synthesis plans with the goal of practicing
green chemistry principles and making intelligent decisions about route selection based on material efficiency in a simple, logical, robust, and reliable manner.

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Appendix 1

**Figure S1**
Convergent steps: step 3, step 7, step 9

Cumulative PMI Expressions:

(i) Main Branch

\[
(cPMI)_{1 \rightarrow 1} = (PMI)_{1} \tag{S1}
\]

\[
(cPMI)_{1 \rightarrow 2} = m_{P2} \left[ (cPMI)_{1 \rightarrow 1} - 1 \right] + (PMI)_{2} \tag{S2}
\]

\[
(cPMI)_{1 \rightarrow 3} = m_{P3} \left[ (cPMI)_{1 \rightarrow 2} - 1 \right] + (PMI)_{3} + \frac{m_{P3}}{m_{P1}} \left[ (cPMI)_{1 \rightarrow 2} - 1 \right] + \frac{m_{P2}}{m_{P1}} \left[ (cPMI)_{1 \rightarrow 2} - 1 \right] \tag{S3}
\]

\[
(cPMI)_{1 \rightarrow 4} = m_{P4} \left[ (cPMI)_{1 \rightarrow 3} - 1 \right] + (PMI)_{4} \tag{S4}
\]

\[
(cPMI)_{1 \rightarrow 5} = m_{P5} \left[ (cPMI)_{1 \rightarrow 4} - 1 \right] + (PMI)_{5} \tag{S5}
\]

\[
(cPMI)_{1 \rightarrow 6} = m_{P6} \left[ (cPMI)_{1 \rightarrow 5} - 1 \right] + (PMI)_{6} \tag{S6}
\]

\[
(cPMI)_{1 \rightarrow 7} = m_{P7} \left[ (cPMI)_{1 \rightarrow 6} - 1 \right] + (PMI)_{7} + \frac{m_{P7}}{m_{P1}} \left[ (cPMI)_{1 \rightarrow 6} - 1 \right] \tag{S7}
\]

\[
(cPMI)_{1 \rightarrow 8} = m_{P8} \left[ (cPMI)_{1 \rightarrow 7} - 1 \right] + (PMI)_{8} \tag{S8}
\]
\[(cPMI)_{1\to g} = \frac{m_B}{m_B} \left[ (cPMI)_{1\to g} - 1 \right] + (PMI)_g + \frac{m_{PMI}}{m_B} \left[ (cPMI)_{g\to 8 \to s} - 1 \right] = (PMI)_{total}\]  

(S9)

(ii) Branch 2

\[(cPMI)_{1*\to 1*} = (PMI)_{1*}\]  

(S10)

\[(cPMI)_{1*\to 2*} = \frac{m_{P2}}{m_{P2}} \left[ (cPMI)_{1*\to 2*} - 1 \right] + (PMI)_{2*}\]  

(S11)

(iii) Branch 3

\[(cPMI)_{2*\to 2*} = (PMI)_{2*}\]  

(S12)

(iv) Branch 4

\[(cPMI)_{4***\to 4***} = (PMI)_{4***}\]  

(S13)

\[(cPMI)_{4***\to 5***} = \frac{m_{P4}}{m_{P4}} \left[ (cPMI)_{4***\to 5***} - 1 \right] + (PMI)_{5***}\]  

(S14)

\[(cPMI)_{4***\to 6***} = \frac{m_{P4}}{m_{P4}} \left[ (cPMI)_{4***\to 6***} - 1 \right] + (PMI)_{6***}\]  

(S15)

(v) Branch 5

\[(cPMI)_{7***\to 7***} = (PMI)_{7***}\]  

(S16)

\[(cPMI)_{7***\to 8***} = \frac{m_{P7}}{m_{P7}} \left[ (cPMI)_{7***\to 8***} - 1 \right] + (PMI)_{8***}\]  

(S17)

Appendix 2

Case scenarios for a two-step plan with respect to sum of step E-factors and overall E-factor.

For a two-step plan given by step 1: \(A + B \rightarrow P_1\), and step 2: \(P_1 + C \rightarrow P_2\) the overall E-factor in terms of the step E-factors \(E_1\) and \(E_2\) is given by equation (S18).

\[E_f = \frac{m_B}{m_{P2}} E_1 + E_2\]  

(S18)

Case I

When \(\frac{m_B}{m_{P2}} = 1\), then \(E_f = E_1 + E_2\).
Case II

When \( \frac{m_{P_1}}{m_{P_2}} > 1 \), then \( E_T > E_1 + E_2 \).

Case III

When \( \frac{m_{P_1}}{m_{P_2}} < 1 \), then \( E_T < E_1 + E_2 \).

Case scenarios for a three-step plan with respect to sum of step E-factors and overall E-factor.

For a three-step plan given by step 1: \( A + B \rightarrow P_1 \), step 2: \( P_1 + C \rightarrow P_2 \), and step 3: \( P_2 + D \rightarrow P_3 \) the overall E-factor in terms of the step E-factors \( E_1, E_2, \) and \( E_3 \) is given by equation (S19).

\[
E_T = \frac{m_{P_1}}{m_{P_2}} E_1 + \frac{m_{P_2}}{m_{P_3}} E_2 + E_3 \quad (S19)
\]

Figure S2 summarizes the 9 possible case scenarios for a three-step plan with respect to the magnitude of the mass ratio of intermediates whether they are equal to 1, greater than 1, or less than 1. Seven of the nine cases yield definite conclusions with respect to the relative magnitudes of the true overall E-factor and the sum of the step E-factors.

**Figure S2**

**Supporting Information:**

Supporting-Information-EXCEL.zip contains calculator-cumulative PMI.xls and six similar files applied to the six synthesis plans for apixaban examined in this work.
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Figure Captions

Figure 1. Target bond mapping for all published synthesis plans to apixaban, 1.
Figure 2. Radial pentagons for Bristol-Myers Squibb 2006 plan for apixaban.
Figure 3. Example reaction network for the synthesis of product P via various synthesis routes from various starting materials showing best available literature yields for each transformation.
Figure 4. Reaction network for syntheses of apixaban.
Figure 5. Flowchart for Stage I of the metrics analyses for documented literature synthesis plans.
Figure 6. Flowchart for Stage II of the metrics analyses for conjectured synthesis plans.
Figure S1. Example Multi-convergent Synthesis Tree Diagram
Figure S2. Summary of overall E-factor versus sum of step E-factors case scenarios for a three-step plan.
STAGE I

Create a reaction network based on known literature documenting all structures of starting materials, intermediate products, and final product; and documenting best reaction yields from the literature for each transformation.

From the reaction network enumerate all possible synthesis routes.

For each route construct a synthesis tree diagram based on balanced chemical equations for each reaction step, labelling MWs of all reagents and intermediate products, stoichiometric coefficients, identifying all branches, labelling reaction yields, and number of steps.

For each synthesis tree diagram use the REACTION and SYNTHESIS spreadsheets to calculate (i) the following step metrics for each reaction: step yield, step AE, step PMI, step E-factor, and step E-factor contributions: E-kernel, E-excess, E-solvent, E-cat, E-workup, and E-purif; construct radial pentagons for each reaction step; and (ii) the following overall metrics for the entire plan: overall yield, overall AE, overall PMI, and overall E-factor contributions.

Construct a metrics results table as shown in Table 1. Rank the synthesis routes according to ascending order of overall PMI; identify best and worst values for the following metrics: overall yield, overall AE, E-kernel, E-excess, E-cat, E-workup, E-purif, E-overall, and overall PMI.

Identify the synthesis route(s) with the most "green" hits in the metrics results table.
STAGE II

Conjecture new synthesis routes to a target molecule structure with a view to obtain a different target bond mapping from published routes.

For each route draw a synthesis tree diagram including all parameters as in Stage I.

Estimate the lowest and highest reaction yields for each step in each route using literature evidence.

For each route calculate the following metrics at the kernel level: overall AE, min. overall yield, max. overall yield, min. kernel overall PMI, max. kernel overall PMI

Rank the synthesis routes according to ascending order of max. overall PMI; identify best and worst values for the following metrics: overall yield and overall AE.

Compare the results of the ranking with those from STAGE I at the kernel level.

Identify any new conjectured routes that show improvements in the kernel metrics.

For each route identify the following drawbacks:
(i) availability of starting materials
(ii) cost of starting materials
(iii) potential for unwanted side reactions
(iv) potential for safety and hazardous materials.

Identify the best candidate conjectured routes satisfying the least drawbacks and the best kernel metrics profile.
\[
\begin{array}{c|c|c}
\frac{m_{R_1}}{m_{R_3}} & \frac{m_{R_2}}{m_{R_3}} & E_T = E_1 + E_2 + E_3 \\
\hline
1 & 1 & E_T = E_1 + E_2 + E_3 \\
1 & > 1 & E_T > E_1 + E_2 + E_3 \\
1 & < 1 & E_T < E_1 + E_2 + E_3 \\
> 1 & 1 & E_T > E_1 + E_2 + E_3 \\
> 1 & > 1 & E_T > E_1 + E_2 + E_3 \\
> 1 & < 1 & \text{indefinite conclusion} \\
< 1 & 1 & E_T < E_1 + E_2 + E_3 \\
< 1 & > 1 & \text{indefinite conclusion} \\
< 1 & < 1 & E_T < E_1 + E_2 + E_3
\end{array}
\]
## Tables

### Table 1. Summary of green metrics for various synthesis plans to apixaban.

| Plan          | # steps | Overall Y (%) | Overall AE (%) | E-kernel | E-excess | E-solvent | E-cat | E-workup | E-purif | E-total | PMI-total |
|---------------|---------|---------------|----------------|----------|----------|-----------|-------|----------|---------|---------|-----------|
| Bristol-Myers | 2006    | 7             | 37.4           | 10.32    | 7.72     | 80.65     | 1.12  | 116.85   | 27.2    | 243.86  | 244.86    |
| Optimus       | 2017    | 9             | 16.7           | 18.4     | 18.63    | 15.90     | 117.40| 2.46     | 214.10  | 9.41    | 377.90    | 378.90    |
| Jiang-Ji      | 2013    | 7             | 26.8           | 18.4     | 11.15    | 16.84     | 104.15| 0.05     | 220.37  | 7.78    | 382.09    | 383.09    |
| MSN           | 2015    | 8             | 16.3           | 19.2     | 24.29    | 28.9      | 126.45| 0.63     | 308.40  | 8.78    | 497.45    | 498.45    |
| Bristol-Myers | 2003    | 7             | 25.6           | 18.5     | 16.06    | 19.15     | 128.22| 0.80     | 460.73  | 9.12    | 624.96    | 625.96    |
| Dr Reddy      | 2017    | 7             | 22.2           | 19.8     | 19.86    | 24.24     | 239.37| 0.86     | 442.27  | 8.78    | 855.96    | 856.96    |

*Calculated along longest branch.

### Table 2. Summary of cumulative complete PMI values for various synthesis plans to apixaban.

| Step j | Optimus-2006 | Jiang-Ji-2013 | Jiang-Ji-2015 | DrReddy-2003 | DrReddy-2017 |
|--------|--------------|---------------|---------------|--------------|--------------|
| 1      | 45.92        | 36.93         | 10.35         | 70.23        | 65.82        |
| 2      | 75.31        | 101.29        | 167.16        | 289.54       |              |
| 3      | 78.81        | 151.55        | 288.44        | 399.80       |              |
| 4      | 95.52        | 214.04        | 354.82        | 579.92       |              |
| 5      | 183.54       | 308.68        | 625.64        | 857.06       |              |
| 6      | 245.01       | 382.83        | 336.47        |              |              |
| 7      | 269.01       |              | 498.35        |              |              |
| 8      | 379.30       |              |              |              |              |
Table 3. Summary of cumulative kernel PMI values for various synthesis plans to apixaban.

| step j | BM-2006 | Optimus-2017 | Jiang-Ji-2013 | MSN-2015 | BM-2003 | DrReddy-2017 |
|--------|---------|--------------|--------------|---------|---------|--------------|
| 1      | 4.30    | 1.79         | 2.68         | 1.71    | 2.59    | 4.00         |
| 2      | 5.69    | 3.45         | 5.36         | 6.48    | 6.06    | 6.28         |
| 3      | 6.11    | 5.64         | 7.59         | 7.78    | 10.06   | 11.43        |
| 4      | 6.20    | 6.97         | 8.07         | 11.99   | 10.87   | 13.86        |
| 5      | 9.90    | 9.88         | 10.29        | 13.64   | 16.56   | 18.65        |
| 6      | 11.17   | 9.19         | 12.05        | 19.79   |         |              |
| 7      | 14.51   |              |              |         |         |              |
| 8      | 19.63   |              |              |         |         |              |

Table 4. Summary of cumulative kernel plus excess PMI values for various synthesis plans to apixaban.

| step j | BM-2006 | Optimus-2017 | Jiang-Ji-2013 | MSN-2015 | BM-2003 | DrReddy-2017 |
|--------|---------|--------------|--------------|---------|---------|--------------|
| 1      | 6.24    | 2.14         | 4.46         | 2.10    | 4.19    | 5.61         |
| 2      | 8.14    | 5.17         | 11.03        | 9.56    | 14.16   | 9.18         |
| 3      | 10.44   | 8.45         | 15.55        | 11.25   | 22.79   | 27.38        |
| 4      | 10.25   | 10.38        | 16.48        | 17.77   | 22.41   | 33.57        |
| 5      | 16.26   | 15.75        | 24.66        | 26.34   | 36.22   | 45.12        |
| 6      | 19.04   | 15.76        | 28.95        | 43.35   |         |              |
| 7      | 25.50   |              |              |         | 54.19   |              |
| 8      |         |              |              |         |         | 35.56        |
| Plan           | complete cPMI | Rank (complete) | kernel cPMI | Rank (kernel) | excess cPMI | Rank (kernel + excess) |
|---------------|---------------|-----------------|-------------|---------------|-------------|------------------------|
| BM-2006       | 245.01        | 1               | 11.17       | 1             | 19.04       | 1                      |
| Optimus-2017  | 379.3         | 2               | 19.63       | 5             | 35.56       | 3                      |
| Jiang-Ji-2013 | 382.83        | 3               | 12.05       | 2             | 28.95       | 2                      |
| MSN-2015      | 498.35        | 4               | 24.73       | 6             | 54.19       | 6                      |
| BM-2003       | 625.64        | 5               | 16.56       | 3             | 36.22       | 4                      |
| DrReddy-2017  | 857.06        | 6               | 18.65       | 4             | 45.12       | 5                      |
### Graphical abstract text:

Key mathematical relationships linking step, overall, and cumulative E-factor and step, overall, and cumulative process mass intensity for any synthesis plan.

**E-factor**

| Step & Overall  | Cumulative          |
|----------------|---------------------|
| $E_{i \rightarrow N} = E_N + \frac{1}{m_{f_N}} \sum_{j=1}^{N-1} m_j E_j$ | $(cE)_{i \rightarrow N} = \frac{m_{f_N}}{m_N} (cE)_{i \rightarrow N-1} + (E)_i$ |

| Process Mass Intensity |
|------------------------|
| $(PMI)_{i \rightarrow N} = (PMI)_N + \frac{1}{m_{f_N}} \sum_{j=1}^{N-1} m_j \left[ (PMI)_j - 1 \right]$ | $(cPMI)_{i \rightarrow N} = \frac{m_{f_N}}{m_N} \left[ (cPMI)_{i \rightarrow N-1} - 1 \right] + (PMI)_i$ |
