DEFT: a program for operators in EFT

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ABSTRACT: We describe a Python-based computer program, DEFT, for manipulating operators in effective field theories (EFTs). In its current incarnation, DEFT can be applied to 4-dimensional, Poincaré invariant theories with gauge group SU(3) × SU(2) × U(1), such as the Standard Model (SM), but a variety of extensions (e.g. to lower dimensions or to an arbitrary product of unitary gauge groups) are possible. Amongst other features, the program is able to: (i) check whether an input list of Lagrangian operators (of a given dimension in the EFT expansion) is a basis for the space of operators contributing to S-matrix elements, once redundancies (such as Fierz-Pauli identities, integration by parts, and equations of motion) are taken into account; (ii) generate such a basis (where possible) from an input algorithm; (iii) carry out a change of basis. We describe applications to the SM (where we carry out a number of non-trivial cross-checks) and extensions thereof, and outline how the program may be of use in precision tests of the SM and in the ongoing search for new physics at the LHC and elsewhere. The code and instructions can be downloaded from http://web.physics.ucsb.edu/~dwsuth/DEFT/.

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1 Introduction

Non-renormalizable quantum field theories, once regarded as something of a pariah by physicists, have become ubiquitous as a means of parameterizing, in a general way, the low-energy effects of unknown physics residing at higher-energy scales. In a nutshell, given a set of quantum fields representing physical degrees of freedom and a group of symmetries acting on them, the lagrangian of such an effective field theory (EFT) contains not just renormalizable invariant operators built out of fields and spacetime derivatives, but all invariant operators, ordered by their relevance in terms of a low-energy expansion. The example that is perhaps of greatest current interest to particle physicists is the use of an EFT given by the renormalizable Standard Model (SM) plus higher-dimension operators (henceforth, the ‘SMEFT’), to parameterize possible deviations from the SM at the Large Hadron Collider and elsewhere.
Whilst the idea of EFT is simple enough in principle, to use it in practice involves a
great deal of donkey work, above and beyond what is required in renormalizable quantum
field theory. There are several reasons for this. Firstly, the operators at a given order in
the low-energy expansion form a vector space whose dimension grows exponentially with
the order. Secondly, given some set of physical observables, there is a large subspace of
operators (whose dimension also grows exponentially with the order of expansion) that
are redundant, in the sense that they do not contribute to any of the observables. These
operators must be identified and dealt with, by forming a basis for the space of physical
observables. Thirdly, in fitting either to experimental data or to some overarching theory,
one must choose a basis for the space of physical operators. Different data sets and different
theories prefer different bases (as do different physicists!) and comparison between them
necessitates a change of basis.

As we shall explain in more detail in section 2, much of the required donkey work
reduces to combinatorics and linear algebra, and is easily done with a computer. To this
end, in this paper we present a computer code, DEFT, to help with the work.

In rough terms, DEFT does the following. At each given order in the EFT dimen-
sion expansion, DEFT generates all possible lagrangian terms that are invariant under the
symmetries of the inputted fields. Through a set of hard coded rules, the program then
enumerates all linear combinations of these operators which do not contribute to S-matrix
elements at said order. By finding the nullspace of these redundant directions, DEFT will
determine a subset of the original lagrangian terms which span the space of physically
distinguishable operators, thereby defining a basis. If given, by the user, any other set of
linear combinations of lagrangian terms, DEFT can check if it too forms a basis. Given two
such bases, DEFT will provide an explicit formula to convert between them. In its current
implementation, DEFT can be applied ‘out-of-the-box’ only to the SMEFT, but the methods
employed may be generalized to a number of phenomenologically relevant EFTs, which we
briefly discuss in section 4.

Experienced practitioners of EFT will easily be able to imagine the benefits of an
automated approach of this type, but let us spell a few of them out anyway.

Firstly, DEFT is able to generate a basis of operators at a given dimension that is not
only (hopefully) correct, but is also obtained relatively quickly, provided that the operator
dimension is not too large. For example, for the SMEFT with one generation of fermions
at operator dimension six, DEFT generates the list of 84 operators in figure 2 in a matter of
minutes. This is to be contrasted with the human approach, which took roughly a quarter
of a century, with more than one hiccough along the way [3, 4].

Secondly, there is a large freedom in the choice of operator basis, which DEFT enables
the user to exploit, according to his or her particular desiderata. There are two aspects
to this freedom. The first corresponds to the usual freedom to choose a basis for a vector
space. But in EFT, there is yet more freedom, which corresponds to the fact that many
operators have the same physical effects. It is often useful, in applications, to exploit this
freedom. On the one hand, for example, an experimentalist whose apparatus is only able to
detect certain types of particles, might prefer a basis description which prioritises operators
containing those particles. On the other hand, an experimentalist whose apparatus detects
only very low energy particles might prefer a basis description with operators containing as few derivatives as possible. Given some input algorithm encoding the user’s desiderata, \texttt{DEFT} will output a corresponding basis. Even, for example, in the SMEFT at dimension 6, the user could simply input a list of 84 operators, and ask \texttt{DEFT} to check that it is a basis. Or the user could input his favourite 10 operators and ask \texttt{DEFT} to generate (if possible) 74 others using its default algorithm or some modification thereof.\footnote{The README file gives some indication as to how bases may be input.}

This freedom to choose a basis has its downsides, of course. Indeed, it seems to be an empirical law of nature that, given an EFT that describes the low-energy limits of some theorists’ models and which is subject to the constraints of some experimentalists’ measurements, the relevant literature is likely to contain roughly as many different choices of basis as the number of theorists and experimentalists put together! This is hardly surprising: the former are likely to choose bases in which the particular operators their theories generate are basis elements and the latter are likely to choose bases in which the operators they constrain best are basis elements. For a few examples of the proliferation of such bases in the SMEFT at $\delta = 6$, the reader is invited to consult, e.g. [5]. As a result, the community has arrived at something of an impasse: in order to compare theory with experiment, or indeed to just compare one experiment with another, phenomenologists must be able to change bases. But such changes of bases are highly non-trivial, because the aforementioned redundancies among operators must be taken into account. Indeed, thus far just one such change of basis has been carried out (by hand) in the SMEFT at dimension 6 [6].

A third, and perhaps the most significant, benefit of \texttt{DEFT} is that such changes of basis can be carried out, not quite at the touch of a button, but with comparable ease. Given an arbitrary set of basis operators, if the user is able to express them in terms of the redundant set of ‘monomial operators’ detailed in section 2.1, \texttt{DEFT} is able to convert them into any other operator basis that the user has so expressed. As an example, we describe the use of \texttt{DEFT} to carry out a change of basis in the SMEFT at dimension 6 in section 3.2. The computation takes 20 minutes on a laptop. We hope therefore, that in removing this impasse \texttt{DEFT} will prove to be useful in the current programme of comparing experimental data with the SM via EFT.

The ability of \texttt{DEFT} to construct arbitrary bases and change between them gives it something of an advantage with respect to recent analytic efforts to determine an EFT basis using Hilbert series methods [7–11]. While these methods are extremely elegant, they naturally require a specific type of basis, namely one in which the numbers of derivatives appearing in operators are minimized. \texttt{DEFT} also enables us to perform an independent cross-check of these methods, and provides an explicit contraction of the Lorentz and gauge indices of the attendant operators.

The genericity of \texttt{DEFT} also distinguishes it from existing Python frameworks with practical applications to (SM)EFT, into which are encoded mappings between particular bases [5], or explicit transformation rules for the conversion between different equivalent operators [12]. For this reason we envisage one use of \texttt{DEFT} to be the construction or conversion between bespoke bases in the Standard Model or similar field theories.
The main drawback of DEFT is that it rapidly runs out of steam as the operator dimension grows. This is hardly surprising, since DEFT works by performing brute-force linear-algebra manipulations in vector spaces whose dimension grows exponentially with the operator dimension, in terms of a redundant description whose size also grows exponentially. Given current computing capabilities, the ceiling corresponds to spaces with roughly $10^3$ physical operators. So in the one-generation SMEFT, going beyond dimension 9 is inconceivable.

The outline of the rest of the paper is as follows. In the next section, we describe the algorithm implemented in DEFT to enumerate operators and redundancies between them, and in section 3 we describe a number of examples and cross-checks, followed by a brief discussion. In appendices A and B respectively, we provide more information on the structure of the code itself, as well as our index conventions. To complement the technical detail of section 2, we provide a more abstract discussion of the structure of the redundancies amongst operators and the related vector spaces in appendices C and D.

2 Implementation

We proceed by describing each step of the program in turn: 1) the construction of a (overcomplete) list of monomial operators (section 2.1); 2) the construction of the IBP (section 2.2), Fierz (section 2.3), commutation of covariant derivatives (section 2.4), and EOM relations (section 2.5); 3) the linear algebra necessary to construct non-redundant bases and to convert into and between them (section 2.6).

2.1 Constructing operators

DEFT assumes that fields transform in irreps of SU($N$), which are described via a combination of upper and lower indices with symmetry conditions attached. An upper index takes values between 1 and $N$ and transforms in the defining rep of SU($N$); a lower index runs between 1 and $N$ and transforms in the conjugate of the defining rep. Conjugation of a field in an irrep of SU($N$) lowers upper indices and vice versa. Presently, DEFT contains the definitions for the fundamental and anti-fundamental irreps, along with the symmetric and traceless combinations thereof.

For our purposes, the irreps of the Lorentz group are those of SU(2)$_{L,lor}$ × SU(2)$_{R,lor}$ — represented by the familiar undotted and dotted indices for the respective SU(2)s of the direct product — with the distinction that, upon conjugation of a field, undotted indices are dotted and vice versa. Table 1 contains the Lorentz and gauge representations, as well as their explicit realisations in terms of (anti)fundamental indices, of the fields of the one generation Standard Model.

The advantage of working with exclusively fundamental and anti-fundamental indices is that there are only two invariant tensors: the Kronecker delta $\delta^a_b$ (with an upper and lower index) and the Levi-Civita epsilon $\epsilon^{abc\ldots z}$ or $\epsilon_{abc\ldots z}$ (with either $N$ upper or $N$ lower indices). We report various sign conventions in appendix B.
Table 1. The fields of the one generation Standard Model in component form, along with their mass dimensions, and their representations under the SM symmetries.

2.2 Integration by parts

Amplitudes which are proportional to a sum of the momenta of the external legs — \( \sum_{i \in \text{external}} k_i^a M_i \) for external leg momenta \( \{k_i^a\} \) — are zero by overall momentum conservation. At the operator level, for each term \( D_\alpha \dot{D}_\beta \ldots D_\gamma \dot{D}_\delta \), we generate a relation by moving the outermost derivative of each field, i.e.,

\[
D_\alpha \dot{D}_\beta \ldots D_\gamma \dot{D}_\delta H = D_\beta \ldots D_\alpha \dot{D}_\delta \dot{D}_\gamma H = \ldots = D_\gamma \ldots D_\delta \dot{D}_\alpha \dot{D}_\beta H = 0.
\]  

(2.1)

\(^2\)A covariant derivative \( D_\alpha \) has a lower SU(2)_{lor} and a lower SU(2)_{lor} index.
2.3 Fierz relations

A product of one upper and one lower Levi-Civita epsilon tensor may be expressed as a sum of products of Kronecker deltas:

\[ \epsilon_{ab\cdots} \epsilon^{xy\cdots} z + \sum_{\xi \in S_N} \sigma(\xi) \delta^x_a \delta^y_b \cdots \delta^z_c = 0, \quad (2.2) \]

summing over the permutations \( \xi \) of the \( N \) upper indices, each having parity \( \sigma(\xi) = \pm 1 \).

For \( N = 2 \), there are also Schouten identities,

\[ \epsilon_{ab} \epsilon_{cd} - \epsilon_{ac} \epsilon_{bd} + \epsilon_{ad} \epsilon_{bc} = 0, \quad (2.3) \]

\[ \delta^a_b \epsilon_{cd} - \delta^c_b \epsilon_{ad} + \delta^{ad}_{bc} = 0, \quad (2.4) \]

\[ \delta^b_a \epsilon_{cd} - \delta^c_d \epsilon_{ad} + \delta^{ad}_{bc} = 0, \quad (2.5) \]

which are effectively ‘raised’ and ‘lowered’ versions of (2.2). \textsc{Deft} searches for the left-hand-most term in each operator, and generates one relation for each match.

In addition, any SU(\( N + k \)); \( k \in \mathbb{N} \) relation of the form (2.2) may have its indices restricted to run between 1 and \( N \), yielding an SU(\( N \)) Fierz relation

\[ \sum_{\xi \in S_{N+k}} \sigma(\xi) \delta^x_a \delta^y_b \cdots \delta^z_c = 0. \quad (2.7) \]

For each set of operators with the same field content having \( N_u \) upper and \( N_l \) lower SU(\( N \)) indices, one such relation is generated for each \( k \leq \min(N_u, N_l) - N \).

2.4 Commuting covariant derivatives

For a field \( G_{xy\cdots}^{ab\cdots} \) which transforms under an SU(\( N \)) gauge group with upper indices \( a, b, \ldots \) and lower indices \( x, y, \ldots \), one can interchange any two of its adjacent covariant derivatives, and the difference of the terms is a sum of field strengths:

\[ (D \cdots D) [D_{\alpha\beta}, D_{\gamma\delta}] (D \cdots D) G_{xy\cdots}^{ab\cdots} = ig_N \left( F_{a\alpha\beta t} G_{xy\cdots}^{ab\cdots} + F_{a\alpha\beta t} G_{xy\cdots}^{ab\cdots} \right. \]

\[ \left. - F_{a\alpha\beta t} G_{xy\cdots}^{ab\cdots} - F_{a\alpha\beta t} G_{xy\cdots}^{ab\cdots} \right). \quad (2.8) \]

where \( g_N \) and \( F_{a\alpha\beta t} ^a = -\frac{1}{2} \epsilon_{\alpha\beta} F_{a\alpha\beta t} ^a - \frac{1}{2} \epsilon_{\alpha\beta} F_{a\alpha\beta t} ^a \) are respectively the gauge coupling and field strengths of the SU(\( N \)) gauge group. One relation is generated per operator per field per adjacent pair of covariant derivatives.

2.5 Equations of motion

The dimension \( n \) part of the following two tree level graphs are equivalent, when all external legs are on-shell and \( n > 4 \): a) a graph comprising a dimension \( n \) vertex and a dimension 4 vertex, and b) a graph comprising a single dimension \( n \) vertex with the same external legs. This is illustrated schematically in figure 1. At the operator level, this corresponds
Figure 1. Two schematic amplitudes whose dimension $n$ parts are equal: the square and circle denote higher and lower derivative dimension $n$ operators.

to redundancies amongst dimension $n$ operators arising from our freedom to make field redefinitions [13]. The redundancies take the form

$$U(x) \frac{\delta S_4}{\delta F(x)} = 0,$$

(2.9)

where $U(x)$ is a functional of some fields, which depend on spacetime coordinate $x$, and $\frac{\delta S_4}{\delta F(x)}$ is an equation of motion (EOM) of the renormalizable theory: the functional derivative of the dimension 4 action w.r.t. a constituent field $F$. Note that $\dim U + 4 - \dim F = n$.

$\text{DEFT}$ constructs the EOM according to the following functional derivative rules:

$$\frac{\delta (DD \ldots D) H(x)}{\delta H(y)} = (DD \ldots D) \delta^{(4)}(x - y),$$

(2.10)

for any (scalar or fermion) matter field $H$;

$$\frac{\delta D_{\alpha\beta} H^a}{\delta A_{\beta\gamma\delta}(x)} = ig_N \delta^a_{\gamma} \delta^\beta_{\delta} \left( \delta^\beta_{\delta} \delta^b_{\gamma} H^b - \frac{1}{N} \delta^d_{\gamma} \delta^a_{\delta} H^a \right) \delta^{(4)}(x - y),$$

(2.11)

for a matter field $H^a$ charged under an SU($N$) gauge group, with coupling $g_N$ and vector potential $A^\gamma_{\alpha\beta}$;

$$\frac{\delta F_{\alpha\beta}}{\delta A^\gamma_{\alpha\beta}(x)} = \epsilon^{\alpha\beta\gamma} \delta^d_{\beta} \left( \delta^\gamma_{\beta} \delta^c_{\gamma} D_{\alpha\delta} - \delta^\gamma_{\delta} \delta^\gamma_{\delta} D_{\beta\gamma} \right) \delta^{(4)}(x - y),$$

(2.12)

for the field strength $F^\alpha_{\alpha\beta}$ of the SU($N$) gauge group, vector potential $A^\gamma_{\alpha\beta}$.

$\text{DEFT}$ also derives the Bianchi identities of the field strengths $F^\alpha_{\alpha\beta}$ by functionally differentiating the action

$$S'_4 = \int d^4x \left( (F^{\alpha\beta})^b_{\alpha\beta}(F_{\alpha\beta})^b_a - (F^{\alpha\beta})^a_{\alpha\beta}(F_{\alpha\beta})^b_a \right),$$

(2.13)

with respect to the corresponding vector potential. Although the Bianchi identities are not EOMs per se, they still give rise to redundancies amongst higher dimensional operators of the form

$$U(x) \frac{\delta S'_4}{\delta A(x)} = 0,$$

(2.14)

and are hence included in this step simply as additional EOMs (one for each gauge group).

Then, for each monomial element of an EOM, the program searches for its embedding in each dimension $n$ term, calculating corresponding ‘quotients’ $U(x)$ (2.9). For each EOM and each possible $U(x)$, relations are formed out of the corresponding terms, weighted by the coefficients of the EOM.
2.6 Reducing to a basis

Having constructed a (overcomplete) list of monomial operators, and a (overcomplete) list of redundancy relations between them, it remains to perform the necessary linear algebra to extract useful bases of operators and the means to convert between them. For this, DEFT uses the Python package sympy [14].

The monomial operators are ordered into a column vector $O_i$, according to a specifiable ranking function. By default, the operators are given a partial ordering to place ‘less desirable’ operators near the top of the vector. Each redundancy relation is then expressed as a row of a matrix $M$, such that $M_{ij} O_j = \mathbf{0}$.

Any valid basis is a set of operators $B_i$ (i.e., a set of linear combinations of $O_i$) which span the nullspace of $M$. DEFT constructs a default basis by placing $M$ in reduced row echelon form (RREF), and defining $B_i$ as the set of monomial operators which do not have a leading entry in any of the rows of the RREF matrix. Combined with the default ordering of $O_i$, this acts to create $B_i$ with as few derivatives as possible (when applied to the Standard Model at dimension 6, for instance, the default $B_i$ closely resemble the operators of the Warsaw basis). The entries of the RREF matrix may be rearranged to express all operators in terms of the basis:

$$O_i = R_{ij} B_j \quad (2.15)$$

The user may specify another putative basis $B'_i$ by manually expressing them in terms of the monomial operators $O_j$, which DEFT encodes as the matrix $B'_i = S_{ij} O_j$. By printing the components of $B'_i = S_{ij} R_{jk} B_k$, DEFT gives expressions to convert between the two bases. The validity of the $B'_i$ as a basis may be checked by the invertibility of the matrix $SR$.

3 Cross checks

3.1 Dimensions

We used the code to calculate the number of independent operators at each mass dimension up to 8, for lagrangians containing various combinations of light fields. The results agree with table 2, whose entries were either computed manually ($d \leq 4$) or using the Hilbert series method of [10] ($d > 4$).\footnote{Operators are compared pairwise by their number of derivatives, then the number of epsilon invariants, then by the number of fields, with the first operator to return a higher number being placed above the other in the column vector. If two operators compare equal by these criteria, and exactly one is (anti)self-conjugate, then said operator is placed lower.}

\footnote{A matrix is in reduced row echelon form if the leading coefficient (the first non-entry from the left) in each row is a 1, each such leading coefficient is the only non-zero entry in its column, and each leading coefficient is to the right of that of the row above it. Its form is invariant under row operations on the original $M_{ij}$.}

\footnote{Both DEFT and table 2 count the dimension 4 operators $F_{\alpha\beta} F^{\alpha\beta}$ and $\tilde{F}_{\alpha\beta} \tilde{F}^{\alpha\beta}$ independently for any field strength $F$.}
Table 2. The number of independent operators at each mass dimension $d$, for various combinations of fields. $\{H, B, W, G, l_L, e_R, q_L, u_R, d_R\}$ are respectively the gauge boson and matter fields of an SU(4) gauge group, with the same electroweak charges as their SU(3) charged counterparts in the Standard Model. In the penultimate line of the table, we treat the SU(4) as a global symmetry.

| $d$ | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
|-----|---|---|---|---|---|---|---|---|
| $\{H, l_L\}$ | 0 | 1 | 0 | 3 | 2 | 6 | 6 | 18 |
| $\{\phi, H, l_L\}$ | 1 | 2 | 2 | 6 | 5 | 12 | 21 | 48 |
| $\{B, e_R\}$ | 0 | 0 | 0 | 3 | 0 | 1 | 0 | 5 |
| $\{H, B, W, l_L, e_R\}$ | 0 | 1 | 0 | 10 | 2 | 23 | 12 | 179 |
| $\{H, B, W, G, l_L, e_R, q_L, u_R, d_R\}$ | 0 | 1 | 0 | 19 | 2 | 84 | 30 | 993 |
| $\{H, B, W, l_L, e_R, q^q_L, u^q_R, d^q_R\}$ | 0 | 1 | 0 | 17 | 2 | 68 | * | * |
| $\{H, B, W, G^4, l_L, e_R, q^q_L, u^q_R, d^q_R\}$ | 0 | 1 | 0 | 19 | 2 | 76 | * | * |

3.2 Change of basis

We define the one generation SILH basis as a one generational restriction of the operators in tables 1, 2, and 3 of [6], and similarly the one generation Warsaw basis from tables 2 and 3 of [4]. DEFT generates expressions for the Wilson coefficients of the SILH basis in terms of the equivalent Wilson coefficients of the Warsaw basis, in agreement with the one generational restriction of the formulae of appendix A of [6], as well as an independent manual calculation.

4 Discussion

In an auxiliary directory in the submission, we provide a list of possible monomial operators in the one generation Standard Model, in the default ordering, together with the reduced row echelon form of $M_{ij}$ when the columns are so ordered.

We depict the structure of the RREF matrix in figure 2, by plotting the non-trivial values of $|R_{ij}|$, as defined in (2.15). Each row of the RREF matrix effectively defines a linear relation expressing each redundant operator in terms of the remaining basis operators; for each row $i$, we plot the absolute values $|R_{ij}|$ on the line of the basis operator corresponding to the index $j$. For each row of the figure, the style of marker is determined by the field composition of the redundant operator ‘being eliminated’, as indicated in the legend.

Note that there are eight lines in figure 2 with no points; equivalently, there are eight columns in the RREF matrix whose entries are all zero, corresponding to six $B$ violating operators and the two operators of the form $H \bar{H} G^2$ and $H \bar{H} \bar{G}^2$. These eight monomial operators are the sole monomial representatives of their respective equivalence classes of $\{ 9 \}$. The operators $O_{Rt}, O_{Rt}, O_{lt}, O_{mu},$ and $O_{uu}$ are absent.

The operators are listed in descending order of number of derivatives, then number of epsilon tensors, indices and conjugate fields.
operators which give the same physical effects in the $S$-matrix, and are, therefore, always in any basis constructed solely from monomial operators. In addition, the remaining two $B$ violating operators — $Q^3L$ and $Q^3\bar{L}$ — are only somewhat trivially related to operators of the same field composition via a Fierz relation. By considering the structure of the redundancy relations, we provide some justification for the apparent isolation of these ten operators in appendix C.

Finally, we would like to consider the potential uses of DEFT beyond that of the Standard Model EFT. In the first step of its procedure, DEFT enumerates all lagrangian terms consistent with the SM symmetries. It does so by representing the fields and derivatives with upper and lower fundamental indices of SU($N$), forming all invariant contractions thereof, and filtering the results for overall zero charge under U(1) symmetries. This step can be performed on an EFT with arbitrary field content, in which the symmetry group is an arbitrary product of unitary groups and whose fields the user is able to encode as irreducible combinations of upper and lower fundamental indices. Other simple conditions, such as invariance under a $Z_2$ symmetry, can similarly be enforced in the filtration step, although adapting DEFT to deal with multidimensional irreps of other symmetry groups would be significantly more involved.

In the second step, the code derives a set of redundancy relations amongst the generated terms — those resulting from integration-by-parts, Fierz identities, replacing commutators of covariant derivatives, Bianchi identities and equation of motion relations, the latter formed using hard-coded rules to functionally differentiate a gauged theory of scalars and fermions. These relations suffice to eliminate all redundancies in the case of the Standard Model EFT, and could be easily reused on another gauge theory with linearly realised symmetries, where these classes of relations are presumed sufficient to eliminate all redundancies in the $S$-matrix. We note, however, that DEFT can provide no guarantee of this sufficiency.

To give one obvious example of a useful extension, given anomalies in data suggesting the need to add light, beyond-the-SM degrees of freedom to the SM, it would be a simple matter to incorporate such fields into DEFT. This has already been done [2] in the case of a SM gauge singlet scalar, hypothesized to explain a spurious anomaly in the $\gamma\gamma$ spectrum at an invariant mass around 750 GeV. Similarly, DEFT could be adapted to use in flavour physics, where the relevant effective lagrangian at the scale of $b$-quarks has SU(3) \times U(1) invariance (where U(1) corresponds to the electromagnetic gauge group) with the Higgs boson, $W, Z$ bosons, and top quark removed. DEFT could also be used to evaluate the restricted set of invariants that arise in theories with some unified symmetry group, such as the SU(4) \times SU(2) \times SU(2) of Pati & Salam or the SU(5) of Georgi & Glashow. With a bit more effort, DEFT could also be applied to theories in lower dimensions, or indeed those with Galileo, rather than Poincaré, invariance.

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Figure 2. The absolute values of the non-trivial components of $R_{ij}$, defined in (2.15), for the one generation SM. Each marker is positioned in line with the basis operator corresponding to index $j$, and formatted according to the field composition of the redundant operator corresponding to index $i$. We define the marginal couplings in terms of the measured coefficients of the three generation Standard Model: the Higgs quartic and gauge couplings equal those measured at the $Z$ pole, whereas the Yukawa couplings are set by measurements of the heaviest generation: $y_u = \frac{m_u}{v}$, $y_d = \frac{m_d}{v}$, $y_e = \frac{m_e}{v}$.
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A Using the code

The code requires Python 2.7+ and the sympy [14] symbolic manipulation package.

Each monomial operator is represented by a Term instance. A Term has a list (.fields) of Field instances and a list (.invariants) of Invariant instances, each of which has a list (.indices) of Index instances that they respectively possess or contract. Field instances also have a list (.Dindices) of tuples of its indices which belong to covariant derivatives acting on the field; a dictionary (.U1Dict) of U(1) charges, which take on rational values represented by frac instances, and a list (.symmetries) of Symmetry instances which enforce symmetry properties of the field’s indices upon contraction with Invariants.

A Relation is a list of Terms (.terms) and corresponding coecients (.weights), which are sympy expressions. sympy is used for some of the subsequent matrix manipulation.

The use of the provided methods for the generation of terms and relations, as well as the conversion into and between bases, is documented in the unit tests, which compute the cross checks described in section 3.

B Index conventions

For an $N$ index group

$$\epsilon^{12\ldots N} = -\epsilon_{12\ldots N} = 1; \quad \delta^a_b = \begin{cases} 1 & \text{if } a = b, \\ 0 & \text{otherwise}. \end{cases}$$ (B.1)

We use the spinor index conventions of [15] with a mostly-minus metric $\eta_{\mu\nu} = \text{diag}(+1,-1,-1,-1)$ and totally antisymmetric tensor $\epsilon^{0123} = -\epsilon_{0123} = +1$. With the

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8The row reduction, when performed symbolically with the marginal couplings of the theory as variables, is computationally expensive. One has the option in DEFT of substituting the different marginal couplings for prime numbers to speed up the row operations, or substituting for zeroes (i.e. working with a free renormalizable part of the theory). Note that one should avoid replacing the couplings with floating point values prior to the row reduction, due to the ensuing propagation of floating point inaccuracies.
use of the tensors $\sigma^{\mu}_{\alpha\dot{\alpha}}$ and $\tilde{\sigma}^{\alpha}_{\mu\dot{\alpha}}$ and relations (2.47)–(2.53) of [15]

$$
\sigma^{\mu}_{\alpha\dot{\alpha}}\tilde{\sigma}^{\beta}_{\mu\dot{\alpha}} = 2\delta^{\beta}_{\dot{\beta}} \delta^{\dot{\alpha}}_{\alpha} ,
$$

(B.2)

$$
\sigma^{\mu}_{\alpha\dot{\alpha}}\sigma_{\mu\beta} = 2\epsilon_{\alpha\beta} \epsilon_{\dot{\alpha}\dot{\beta}} ,
$$

(B.3)

$$
\tilde{\sigma}^{\alpha\alpha} \tilde{\sigma}^{\beta\beta} = 2\epsilon^{\alpha\beta} \epsilon^{\dot{\alpha}\dot{\beta}} ,
$$

(B.4)

$$
[\sigma^{\mu\dot{\nu}} + \sigma^{\nu\dot{\mu}}]_{\alpha\dot{\beta}} = 2\eta^{\mu\nu} \delta^{\alpha\dot{\beta}} ,
$$

(B.5)

$$
[\tilde{\sigma}^{\mu\dot{\nu}} + \tilde{\sigma}^{\nu\dot{\mu}}]_{\dot{\alpha}\alpha} = 2\eta^{\mu\nu} \delta_{\dot{\alpha}\alpha} ,
$$

(B.6)

$$
\sigma^{\mu\dot{\nu}} \sigma^\nu = \eta^{\mu\nu} \sigma^\nu - \eta^{\nu\mu} \sigma^\mu + \eta^{\nu\nu} \sigma^\mu - i\epsilon^{\mu\nu\rho\kappa} \sigma^\kappa ,
$$

(B.7)

$$
\tilde{\sigma}^{\mu\dot{\nu}} \tilde{\sigma}^\nu = \eta^{\mu\nu} \tilde{\sigma}^\nu - \eta^{\nu\mu} \tilde{\sigma}^\mu + \eta^{\nu\nu} \tilde{\sigma}^\mu - i\epsilon^{\mu\nu\rho\kappa} \tilde{\sigma}^\kappa ,
$$

(B.8)

expressions involving vector and spinor Lorentz indices may be easily converted. For expressions with a single vector index we define

$$
A_{a\dot{a}} = \sigma^{\mu}_{a\dot{a}} A_{\mu} ,
\quad D_{a\dot{a}} = \sigma^{\mu}_{a\dot{a}} D_{\mu} ,
$$

(B.9)

whence we derive

$$
A_{\mu} = \frac{1}{2} \tilde{\sigma}^{\alpha\dot{\alpha}} A_{a\dot{a}} ,
\quad D_{\mu} = \frac{1}{2} \tilde{\sigma}^{\dot{\alpha}\alpha} D_{a\dot{a}} .
$$

(B.10)

For expressions with two vector indices, such as a field strength $F_{\mu\nu}$ or its dual $\tilde{F}_{\mu\nu}$, we define

$$
F_{a\dot{a}b\dot{b}} = \sigma^{\mu}_{a\dot{a}} \sigma^{\nu}_{b\dot{b}} F_{\mu\nu} ,
\quad \tilde{F}_{a\dot{a}b\dot{b}} = \sigma^{\mu}_{a\dot{a}} \sigma^{\nu}_{b\dot{b}} \tilde{F}_{\mu\nu} ,
$$

(B.11)

such that

$$
F_{\mu\nu} = \frac{1}{4} \tilde{\sigma}^{\alpha\dot{\alpha}} \sigma^{\beta\dot{\beta}} F_{a\dot{a}b\dot{b}} ,
\quad \tilde{F}_{\mu\nu} = \frac{1}{4} \tilde{\sigma}^{\dot{\alpha}\alpha} \sigma^{\dot{\beta}\beta} \tilde{F}_{a\dot{a}b\dot{b}} ,
$$

(B.12)

where $F_{a\dot{a}b\dot{b}}$ and $\tilde{F}_{a\dot{a}b\dot{b}}$ may be expressed in terms of Lorentz group irreps $F_{\alpha\beta}$ and $\tilde{F}_{\dot{\alpha}\dot{\beta}}$:

$$
F_{a\dot{a}b\dot{b}} = -\frac{1}{2} (\epsilon_{\dot{a}\beta} F_{a\dot{a}b\dot{b}} + \epsilon_{a\dot{b}} F_{\dot{a}a\dot{b}b}) ,
\quad \tilde{F}_{a\dot{a}b\dot{b}} = -\frac{1}{2} (\epsilon_{\dot{a}\beta} F_{\dot{a}a\dot{b}b} - \epsilon_{a\dot{b}} F_{a\dot{a}b\dot{b}}) .
$$

(B.13)

If $F_{\mu\nu} = \partial_{\mu} A_{\nu} - \partial_{\nu} A_{\mu}$, helpful consequences of the above conventions include

$$
F_{\alpha\beta} = \epsilon^{\dot{\alpha}\dot{\beta}} F_{\dot{\alpha}a\dot{\beta}b} = \epsilon^{\dot{\alpha}\dot{\beta}} (\partial_{a\dot{a}} A_{\beta\dot{b}} - \partial_{\beta\dot{b}} A_{a\dot{a}}) ,
$$

(B.14)

$$
\tilde{F}_{\dot{\alpha}\dot{\beta}} = \epsilon^{\alpha\beta} F_{\dot{\alpha}a\dot{\beta}b} = \epsilon^{\alpha\beta} (\partial_{\alpha\dot{a}} A_{\dot{\beta}\dot{b}} - \partial_{\dot{\beta}\dot{b}} A_{\alpha\dot{a}}) .
$$

(B.15)

Note that, alternatively, one can use the tensors

$$
(\sigma^{\mu\nu})_{\alpha}^{\beta} = \frac{1}{4} i(\sigma^{\mu\nu}_{\alpha\dot{a}} \sigma^{\nu\alpha}_{\dot{a}\beta} - \sigma^{\nu\nu}_{\alpha\dot{a}} \sigma^{\mu\alpha}_{\dot{a}\beta})
$$

(B.16)

$$
(\tilde{\sigma}^{\mu\nu})_{\dot{\alpha}}^{\dot{\beta}} = \frac{1}{4} i(\tilde{\sigma}^{\mu\nu}_{\alpha\dot{a}} \sigma^{\nu\alpha}_{\dot{a}\beta} - \tilde{\sigma}^{\nu\nu}_{\alpha\dot{a}} \sigma^{\mu\alpha}_{\dot{a}\beta})
$$

(B.17)

to convert directly between different forms of the field strength:

$$
F_{\alpha\beta} = 2i(\sigma^{\mu\nu})_{\alpha}^{\beta} \epsilon_{\gamma\beta} F_{\mu\nu} ;
\quad \tilde{F}_{\dot{\alpha}\dot{\beta}} = 2i(\tilde{\sigma}^{\mu\nu})_{\dot{\alpha}}^{\dot{\beta}} \tilde{F}_{\mu\nu} ;
$$

(B.18)

$$
F_{\mu\nu} - i\tilde{F}_{\mu\nu} = \frac{1}{2} i F_{\alpha\gamma} \epsilon^{\gamma\beta} (\sigma^{\mu\nu})_{\alpha}^{\beta} ;
\quad F_{\mu\nu} + i\tilde{F}_{\mu\nu} = \frac{1}{2} i \epsilon^{\gamma\beta} \tilde{F}_{\beta\dot{\gamma}} (\sigma^{\mu\nu})_{\dot{\alpha}}^{\dot{\beta}} .
$$

(B.19)
A four component Dirac spinor may be expanded in terms of the components of a left-handed, $L^\alpha$, and right-handed, $R^\alpha$, Weyl spinor, such that

$$\Psi = \begin{pmatrix} L_1 \\ L_2 \\ R^1 \\ R^2 \end{pmatrix}$$

and its conjugates are

$$\bar{\Psi} = \begin{pmatrix} \bar{L}^\alpha \\ \bar{L}_\dot{\alpha} \end{pmatrix}; \quad \bar{\Psi}^C = \begin{pmatrix} \bar{R}_\alpha \\ \bar{R}_{\dot{\alpha}} \end{pmatrix}.$$  

Gamma matrices may be similarly expanded as

$$\gamma^\mu = \begin{pmatrix} 0 & \sigma^\mu_{\alpha\beta} \\ \sigma^{\mu\dot{\alpha}\dot{\beta}} & 0 \end{pmatrix}; \quad \gamma^5 = \begin{pmatrix} -\delta^\alpha_\alpha & 0 \\ 0 & \delta^\dot{\alpha}\dot{\alpha} \end{pmatrix}; \quad \frac{1}{4} i [\gamma^\mu, \gamma^\nu] = \begin{pmatrix} (\sigma^{\mu\nu})^\alpha_{\dot{\alpha}} & 0 \\ 0 & (\sigma^{\mu\nu})^\dot{\alpha}\alpha \end{pmatrix}.$$  

We normalize the non-Abelian vector potentials of the SM such that

$$(W_{a\dot{a}})^b = \frac{1}{2} W^i_{a\dot{a}} (\sigma^i)^a_{\dot{a}}; \quad (G_{a\dot{a}})^b = \frac{1}{2} G^i_{a\dot{a}} (\lambda^i)^a_{\dot{a}},$$

where $(\lambda^i)^a_{\dot{a}}$ is the value of the $a$th row and $\dot{a}$th column of the $i$th Gellmann matrix, and similarly for the Pauli sigma matrices $(\sigma^i)^a_{\dot{a}}$. $G^i, i = 1, \ldots, 8,$ and $W^i, i = 1, \ldots, 3,$ are the canonical gauge fields found in, for instance, the listing of the Warsaw basis [4]. With the use of the Fierz relations,

$$(\sigma^i)^a_{\dot{a}} (\sigma^j)^b_{\dot{b}} = 2 \delta^a_{\dot{a}} \delta^b_{\dot{b}} - \delta^a_{\dot{b}} \delta^b_{\dot{a}},$$

$$(\lambda^i)^a_{\dot{a}} (\lambda^j)^b_{\dot{b}} = 2 \delta^a_{\dot{a}} \delta^b_{\dot{b}} - \frac{2}{3} \delta^a_{\dot{b}} \delta^b_{\dot{a}},$$

we can deduce the correct normalization of the kinetic terms, e.g.,

$$-\frac{1}{4} W^i_{\mu\nu} W^{i\mu\nu} = \frac{1}{16} \left( (W^a_{\alpha\beta})_b^a (W_{a\alpha\beta})_d^b + (W^{a\dot{\alpha}\dot{\beta}})_b^{a\dot{\alpha}\dot{\beta}} (W_{a\dot{\alpha}\dot{\beta}})_d^b \right),$$

$$-\frac{1}{4} G^i_{\mu\nu} G^{i\mu\nu} = \frac{1}{16} \left( (G^{a\dot{\alpha}\dot{\beta}})_b^{a\dot{\alpha}\dot{\beta}} (G_{a\dot{\alpha}\dot{\beta}})_d^b + (G^{a\dot{\alpha}\dot{\beta}})_b^{a\dot{\alpha}\dot{\beta}} (G_{a\dot{\alpha}\dot{\beta}})_d^b \right).$$

C The structure of operator relations in a generic 4d EFT

Following the procedure of [16], we define two integer coordinates $n$ and $\sum h$ for each monomial EFT operator as, respectively, the number of fields and the sum of the helicities $h$ of the particle created by the action of each field on the vacuum. For fields that are scalar $\phi$, left- and right-handed Weyl fermions $\psi$ and $\bar{\psi}$, and left- and right-handed field strengths $F$ and $\bar{F}$, $h = 0, +\frac{1}{2}, -\frac{1}{2}, +1, -1$ respectively. We enumerate the possible field compositions of dimension 6 operators allowed by Lorentz symmetry and arrange them by their coordinates in figure 3 (cf. figure 1 of [16]).

---

9For the purposes of calculating $n$ and $\sum h$, we treat covariant derivatives as partial derivatives.
Consider how redundancy relations allow one to move around the table of figure 3. IBP and Fierz relations ‘trivially’ mix operators with the same field composition, and therefore with the same coordinates \((n, \sum h)\). The other two kinds can be viewed as expressing a higher derivative operator in terms of an equivalent sum of lower derivative ones.

One, replacing a commutator of derivatives with a field strength generically yields a combination of terms, some with an additional \(F\), some with an \(\tilde{F}\) (one of these may be forbidden by Lorentz symmetry). Thus, starting with an operator with coordinates \((n, \sum h)\), one ends up with operators at \((n + 1, \sum h + 1)\) and \((n + 1, \sum h - 1)\).

Two, replacing the free part of an EOM with the interacting parts amounts to, diagrammatically, taking a graph comprising just an insertion of a higher derivative dimension 6 operator, and adding a dim 4 vertex to one of the legs on which the derivative(s) act(s) (see figure 1). This composite, two vertex graph may have the same leading order momentum piece as a simple insertion of a lower derivative dim 6 operator, when the external legs are on shell. We may assume the fields are massless, as relevant interactions do not affect the EOM relations.\(^\text{10}\) Thus, by (12) of [16], the coordinates \((n_i, (\sum h)_i)\) of such a composite amplitude (and by extension the weights of the corresponding lower derivative operator) are related to the weights of the constituent vertices \((n_j, (\sum h)_j)\) and \((n_k, (\sum h)_k)\) by:

\[
\begin{align*}
n_i &= n_j + n_k - 2; \\
\left(\sum h\right)_i &= \left(\sum h\right)_j + \left(\sum h\right)_k.
\end{align*}
\]

(C.1)

The weights of possible dim 4 vertices are as follows. A gauge or Yukawa coupling is \((n, \sum h) = (3, \pm 1)\). Anything proportional to a scalar quartic is \((4, 0)\). Therefore, the part of an EOM relation proportional to a gauge or Yukawa coupling lies one unit right and one unit either up or down in the table of operators, relative to the original higher derivative operator. For the part proportional to a Higgs quartic, it lies two units to the right.

Figure 3 allows us to understand two examples of dimension 6 monomial operators in the SM, which are not related to any others. One, an \(H^2G^2\) (class \(F^2\phi^2\)) operator could only be reached from an operator of class \(F\phi^2D^2\). However, all such \(GH^2D^2\) operators are forbidden by gauge symmetries. Two, baryon violating operators of the form \(\psi^4\), \(\psi^2\psi^2\), and \(\psi^4\), are only reachable from operators of class \(F\psi\tilde{\psi}D\) and \(\psi^2D^3\), as well as their conjugates. The baryon violating operators contain three quarks, and all relations preserve the parity of the number of quarks. However, there are no gauge invariant operators of the form \(F\psi\tilde{\psi}D\) or \(\psi^2D^3\) containing a single quark, leaving the baryon violating four fermion operators unrelated to both baryon conserving operators, and also unrelated to each other.

D Vector spaces of operators and observables

Though ultimately all of the computations \textsc{DEFT} carries out are performed in a specific basis or bases, we find it illuminating to consider its operations abstractly in a way that is basis independent.

We thus define the operators of a given dimension as the gauge and Poincaré invariants built out of formal combinations of fields and spacetime derivatives. We insist that

\(^{10}\)More precisely, the effects of mass terms in the EOMs can be absorbed by redefinitions of the dim 4 coefficients in the lagrangian.
Figure 3. A schematic ‘map’ of all dimension 6 operators allowed by Lorentz symmetry, cut in half about its axis of symmetry $\sum h = 0$ (reflected in this line are the hermitian conjugates of the operators shown). Arrows show the movement induced by equation of motion relations and commutation of derivative relations in the space of dim 6 operators, colour-coded to show $\phi$ EOMs (blue), $\psi$ EOMs (green, short dashed), $F$ EOMs (red, long dashed), and replacing derivatives with field strengths (grey, dash-dotted).

these be hermitian.\(^{11}\) Since the sum of two such operators is itself an operator, and since multiplying an operator by a real number yields an operator, \&c, the operators form a (finite-dimensional) vector space, $V$ over the reals.

At a given order in the expansion, the observables may be regarded as maps from $V$ to $\mathbb{R}$, where the value of the map is given by the real number that would be obtained by a measurement of the observable, given the theory corresponding to that operator. At the given order, the operator contributes to the observable via interference with lower dimension operators, and so the map is linear. Hence the observables are elements of the dual vector space $V^*$.

Not every element of $V^*$ (not every linear map) can be an observable, however. For one thing, the operators in $V$, which we regard as formal combinations of fields and derivatives, may be subject to underlying mathematical identities. For example, some linear combination of operators may satisfy a Fierz or Schouten identity, or be a total derivative. As a result (at least in perturbation theory), all observables must yield zero on those linear combinations of formal operators. Moreover, at least in particle physics collider experiments

\(^{11}\)We remark that, as is common in the literature, DEFT deals with non-hermitian operators with correspondingly complex coefficients. Since this confuses the counting of operators, we insist that they be hermitian in this appendix, such that all the resulting vector spaces are isomorphic to $\mathbb{R}^n$. 

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(although not necessarily in other areas of physics), observables are restricted to $S$-matrix observables (things which can be measured ‘at infinity’) and one may show \cite{13} that such observables yield zero for any operator that vanishes when the renormalizable equations of motion hold, up to corrections that are higher order in momentum counting. Finally, it may happen that, given our current technological limitations, some things are simply not observable, or that we are simply not interested in them. Thus it is useful to define a subspace $U^* \subset V^*$ of observables of interest.

Given $U^*$, it is natural to consider the space $W \subset V$ of redundant operators defined as the operators that yield zero for all observables. It then follows that $U \cong (U^*)^*$ is simply given by $V - W$, the quotient space obtained by identifying any two operators in $V$ that differ by an operator in $W$. $U$ is also a vector space (though it is not a subspace of $V$!) and we call it the space of physical operators. We stress (as in \cite{1}) that the elements of $U$ are equivalence classes of operators in $V$, where the equivalence relation is defined such that any two operators in $V$ whose difference lies in $W$ are considered equivalent.

We stress again that, according to our definition, $U^*$ excludes not only observables that are ‘mathematically unobservable’, in the sense of being related by underlying identities that hold irrespective of what we do and do not observe, but also those that are unobservable because of the restricted nature of the experiments that we have in mind. We find this to be a useful concept, as the following examples illustrate.

Suppose, for example, that, much like the ancient Greeks, our experiments are purely of the *gedanken* variety, such that we don’t bother to measure anything. Then $U^* = \{0\}$ contains only the zero vector, $W = V$, and $U = \{0\}$, such that there are no non-trivial physical operators.

Alternatively, suppose that we are only interested in searching for baryon number violation by $\pm 1$ units at dimension 6 in the SMEFT, and so restrict our attention to experiments sensitive to processes in which baryon number is violated by $\pm 1$ units. Then $W$ contains all operators in which baryon number is violated by some other number of units, because at this order, such operators can only interfere with lower dimension processes in the SMEFT, all of which conserve baryon number. The physical operators in $U$ are then those operators which violate baryon number by $\pm 1$ units.\textsuperscript{12}

Finally, suppose that we build a ‘superdupercollider’ in which all $S$-matrix elements are observable. The corresponding $U$, which is parametrised by the bases in DEFT, contains all operators that do not vanish under mathematical identities or when the equations of motion hold.

\textsuperscript{12}This example admits the following generalization. At a given dimension, we may consider a symmetry of the lagrangian at lower dimensions, accidental or otherwise, and reduce operators into combinations carrying real irreducible representations of that symmetry. (The representations are real because the operators in the lagrangian are elements of a real vector space. Hence the need to consider processes violating baryon number by either $+1$ or $-1$ units in the example.) Any collection of these irreps can be associated with a corresponding subspace of observables in a similar way.
D.1 Remarks on truncations

Since the dimension of the space of physical operators grows exponentially with the order of the EFT expansion, it tends to quickly become unmanageably large. For example, in the SMEFT with 1 generation of fermions, the space is 1 dimensional in $d = 0$ and 2 (corresponding to the vacuum energy and Higgs mass parameter, respectively, and 19, 84, and 993 dimensional in $d = 4, 6, 8$ respectively [4, 10].

Given this state of affairs, it is natural to try to reduce the dimension of the space by some kind of truncation. We have already shown how this can be done by restricting to the space of physical observables of interest and defining a corresponding space of redundant operators as those which do not contribute to the observables of interest. The space of physical operators is then obtained as the quotient space.

Many authors have gone further, in restricting to a subspace of operators on the basis of some kind of theoretical prejudice. Though it is somewhat out of the main thrust of this paper, we feel it worthwhile to issue some parenthetical remarks regarding the pitfalls of such an approach.

To be concrete, the typical strategy is to pick a ‘theoretically-motivated’ list of operators and then to consider just the span of those operators in $V$ in fits to data. Now, it is certainly the case that such a span defines a subspace of $V$ and, correspondingly, a subspace of the space of physical operators $U$. Each vector in the latter subspace represents a perfectly viable theory (at least from the EFT perspective) and so one may sensibly ask whether the data rule it out or not. But one should be very careful in trying to assign some physical meaning to the span of the operators as a whole. Indeed, such a meaning can only be unambiguous if it is well-defined on $U$, i.e., on the equivalence classes in $V$.

It is perhaps easiest to illustrate the danger by means of an explicit example. Suppose, for example, (as has been done in the literature) that one is interested in the possibility of new physics effects in the top quark sector. Given a basis of operators for $V$, one could then try to truncate by retaining only operators featuring a top quark field in that basis. But such a truncation certainly does not correspond to the class of theories with new physics in the top sector, because it is not well-defined on the equivalence classes of physical operators! Indeed, it is a choice which depends arbitrarily on the basis that we choose for $V$. If we change to a basis in which we replace an operator involving a top quark with an operator not involving a top quark, then the truncated space of physical operators that we obtain will also change.

For another example, suppose that we try to divide operators into the order at which they can be generated in a renormalizable UV completion. So, for example, we might consider only the operators that can be generated at tree-level. But the meaning of this is ambiguous, because it is not, in general, well-defined on the equivalence classes. Indeed, a number of counterexamples in the SMEFT are given in [1].

These ambiguities can be avoided by truncating directly on the equivalence classes themselves. The problem, of course, is that the equivalence classes are rather difficult to characterise. DEFT can be used to help with the characterization. As an example of this, in

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[13] The number is reduced to 17 if one eliminates the operators $B\bar{B}$ and $W\bar{W}$; these are retained in DEFT.
section 3 we provide a spanning set of unobservable directions in the SMEFT and describe some of their properties.

We have already given one example of a manifestly consistent truncation, namely in dividing operators into the real, irreducible representations they carry of the symmetries (accidental or otherwise) of the lower-order lagrangian. Thus, in the SMEFT at \( d = 5 \) we may classify the operators according to the representations of baryon and lepton family numbers that they carry, while at \( d = 6 \) we may classify them by their baryon number and lepton parity; for \( d \geq 7 \) no accidental symmetries remain.

Many other possible truncations remain. Indeed, any truncation of the space of physical operators will do. But, presumably, some of these truncations are more natural than others. For example, an inspection of the redundancy relations shows that the \( d = 6 \) operator containing fields \( GGHJI \) is in a class of its own, such that it always appears in a basis. We provide an argument for this based on the general structure of operator redundancies described in appendix C.

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