Fermion RG blocking transformations and IR structure

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We explore fermion RG block-spinning transformations on the lattice with the aim of studying the IR structure of gauge theories and, in particular, the existence of IR fixed points for varying fermion content. In the case of light fermions the main concern and difficulty is ensuring locality of any adopted blocking scheme. We discuss the problem of constructing a local blocked fermion action in the background of arbitrary gauge fields. We then discuss the carrying out of accompanying gauge field blocking. In the presence of the blocked fermions implementation of MCRG is not straightforward. By adopting judicious approximations we arrive at an easily implementable approximate RG recursion scheme that allows quick, inexpensive estimates of the location of conformal windows for various groups and fermion representations. We apply this scheme to locate the conformal windows in the case of SU(2) and SU(3) gauge groups. Some of the reasons for the apparent efficacy of this and similar decimation schemes are discussed.

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1. Introduction

The dependence of the phase structure of non-Abelian gauge theories on their fermion content is a question of obvious field theoretic and physical interest. In view of anticipated discoveries at the LHC this question has recently come to the fore with some urgency. For sufficiently small number of fermions there is UV asymptotic freedom (AF) and IR confinement. It is also known that for sufficiently large number of fermions asymptotic freedom and confinement are lost. It is generally believed that there is an intermediate range with an UV AF fixed point but no confinement; instead there should be an IR fixed point resulting in conformal IR dynamics. The dependence of these ‘conformal windows’ on the gauge group, the number of fermion multiplets and their representation, as well as the structure of the associated IR fixed points, i.e. their relevant directions and anomalous dimensions, are questions of central importance for informed model building "beyond the standard model". For example, the viability of current walking technicolor models depends on such detailed knowledge.

Several approaches have been employed over the last few years in order to investigate these issues (see [1] - [2] for review and references). One popular method is to adopt some definition of a running coupling, e.g. the ‘Schroedinger functional’ definition. Since any definition of a ‘coupling’ outside the perturbative regime is necessarily arbitrary, any such choice is scheme-dependent. Caution should then be exercised in interpreting results. Another approach is to compute the spectrum and/or some judiciously chosen physical observable and track their deformation as a function of the number of fermions and their masses. Such spectral studies should, in principle, yield a clear outcome, but they are very expensive to carry out in practice to the degree of accuracy required for unambiguous results. Furthermore, it is not always clear what the expected behavior should be.

A direct method for getting at the phase structure would be implementation of the Wilsonian Renormalization Group (RG). Successive RG blocking transformations determine the effective action RG flow and its fixed points. Conceptually, this is the most straightforward and unambiguous approach. Various RG-based studies have been carried out recently for the most part using the two-lattice matching MCRG method [4] - [7]. There are, however, special challenges that must be met when attempting to implement RG blockings and MCRG methods in the presence of light fermions.

In the following we will examine some of the issues involved in performing RG transformations with fermions. We will examine the question of preserving locality of the resulting action after an RG blocking step of light fermion degrees of freedom. For full RG implementation such a step must of course be accompanied by a gauge field blocking step. Again, how to do this in practice in the presence of fermions, in particular how to implement MCRG, is not straightforward. To get some insight we will adopt some judicious approximations that lead to an easily implementable approximate RG recursion scheme [8]. This scheme, which proves rather effective, is then applied to locating the conformal windows in the case of the $SU(2)$ and $SU(3)$ groups. Some or the reasons for its apparent effectiveness are discussed in the last section.

2. Fermion RG blocking

In devising RG blocking transformations for light fermions the main concern is maintaining
reasonable locality of the blocked action after each blocking step. This is a question of interest in it’s own right apart from issues of practical implementation. For free fermions a blocked action can be defined along the lines below (eq. (2.3)-(2.4)) that can be shown to maintain locality [9]. This has been used to argue [10] the locality of determinant rooting for free fermions. This locality demonstration, however, cannot be extended in the presence of an arbitrary gauge field background.

To begin exploring this issue consider the Wilson operator in background gauge field U:

\[ D_0(U) = 1 - \kappa M[U], \]

(2.1)

where the only non-vanishing elements of matrix M are:

\[ M_n (n+\mu)[U] = (1 - \gamma_\mu) U_\mu(n), \quad M_n (n-\mu)[U] = (1 + \gamma_\mu) U_\mu^*(n-\mu). \]

(2.2)

A fermion RG blocking step specifies a transition from the fermion degrees of freedom on the original lattice \( \Lambda \) to ‘thinned-out’ degrees of freedom on a lattice \( \Lambda' \). With \( \Lambda \) a lattice of spacing \( a \), let \( \Lambda^{(1)} \) denote the lattice of spacing \( 2a \). We take \( \Lambda' = \Lambda^{(1)} \cup S \), where \( S \) is a subset of \( \Lambda \) to be specified later.

We introduce variables \( \Psi^{(i)} \) on \( \Lambda' = \Lambda^{(1)} \cup S \) by

\[ \int D\bar{\Psi} D\Psi \exp(-\bar{\Psi} D_0 \Psi) = \int D\bar{\Psi} D\Psi D\bar{\Psi}^{(i)} D\Psi^{(i)} \exp \left\{ -\bar{\Psi} D_0 \Psi - \alpha (\bar{\Psi}^{(i)} - \Psi^{(i)}) (\Psi^{(i)} - Q \Psi) \right\}. \]

(2.3)

The rectangular matrices \( Q : \Lambda \to \Lambda' \) are given by

\[ Q_{nm} \Psi_m = \Psi_n + \zeta \sum_{\mu=1}^d M_n (n+\mu)[U] \Psi_n(n+\mu) + \zeta \sum_{\mu=1}^d M_n (n-\mu)[U] \Psi_n(n-\mu) \]

(2.4)

\[ = \left( \delta_{nm} + \zeta \tilde{Q}_{nm} \right) \Psi_m. \]

(2.5)

Note that \( \alpha \to \infty \) gives a \( \delta \)-function definition of the \( \Psi^{(i)} \)'s, i.e. \( \Psi^{(i)}_n = (Q \Psi)_n \). Integrating out the original variables one has on the new lattice \( \Lambda' \):

\[ \int_\Lambda D\bar{\Psi} D\Psi \exp(-\bar{\Psi} D_0 \Psi) = \text{Det}^{-1} \int_{\Lambda'} D\bar{\Psi}^{(i)} D\Psi^{(i)} \exp \left\{ -\alpha \bar{\Psi}^{(i)} \Psi^{(i)} + \alpha^2 \bar{\Psi}^{(i)} Q G \Psi^{(i)} \right\} \]

(2.6)

with

\[ G^{-1} = \left[ D_0 + \alpha Q^* \right]. \]

(2.7)

In general, with light fermions, \( G \) is a very non-local propagator.

We now choose the set \( S \) and the decimation parameter \( \zeta \) in (2.4) so that in the action

\[ -\bar{\Psi} D_0 \Psi - \alpha (\bar{\Psi}^{(i)} - \Psi^{(i)}) (\Psi^{(i)} - Q \Psi) \]

in (2.3) one cancels the nontrivial part of \( D_0 \):

\[ \bar{\Psi} \left[ \kappa M - \alpha \zeta (\tilde{Q} + \tilde{Q}^*) \right] \Psi = 0. \]

(2.8)
This is accomplished by setting $\zeta = \kappa / \alpha$, and taking the set $S$ as shown in Fig. 1(a), i.e. $S$, viewed as a subset of $\Lambda$, is the set of the interior sites of the boundary plaquettes of the 2x2-hypercubes that are the elementary d-cells of $\Lambda'$. The propagator $G$ for the the original variables $\psi$ in (2.3) is then given by

$$G^{-1} = 1(\alpha) + \frac{\kappa^2}{\alpha} \tilde{Q}^\dagger \tilde{Q} > 0, \quad 1(\alpha)_{nm} = \begin{cases} \frac{1 + \alpha}{\delta_{nm}} & n, m \in \Lambda' \\ \delta_{nm} & n, m \in \Lambda \setminus \Lambda' \end{cases},$$

and the action in (2.6) from the integration over the original $\psi$ becomes

$$\bar{\Psi}^{(1)} \left[ \frac{\alpha}{1 + \alpha} - \kappa^2 \tilde{Q} G \tilde{Q}^\dagger \right] \Psi^{(1)}.$$  \hspace{1cm} (2.10)

(2.10) is manifestly local on $\Lambda' = \Lambda^{(1)} \cup S$. In particular, note that from (2.9), taking $\alpha$ large, interactions induced by $G$ may be expanded in a hopping expansion in $\frac{\kappa^2}{\alpha}$ and resummed. In fact, in the limit $\alpha \to \infty$, i.e. for the $\delta$-function definition of the blocked variables (cf. above), $G$ becomes ultralocal, and in fact $U$-independent, and the action goes to

$$\bar{\Psi}^{(1)} \left[ 1 - \kappa^2 \tilde{Q} \tilde{Q}^\dagger \right] \Psi^{(1)}.$$  \hspace{1cm} (2.11)

After this blocking step we end up with fermions on the boundary of the blocked cube, i.e. a form of exact ‘potential moving’ fermion blocking albeit on a non-hypercubic lattice. The fermion action is still bilinear in the blocked fermion fields which interact via ‘fat’ gauge field bonds (Fig. 1(b)). The gauge field bond variables are yet to be blocked.

It is important to note that if one starts integrating out gauge field bonds inside each blocked hypercube, four-fermion and higher-fermion interactions will in general be induced (but of course at the scale of the blocked cube). Ultralocality of $G$ in this connection completely avoids potential difficulties from the $\text{Det}G^{-1}$ factor in (2.6) in performing any such accompanying gauge field blocking.

We have devised a fermion blocking step in a general gauge field background that results in a manifestly local action. How to perform successive fermion blocking steps in a fixed gauge field
background under the requirement that the action be manifestly local at the scale of each step is an interesting question in its own right. The following is a sufficient condition: for the n-th step implement the above blocking procedure integrating out the fermions on the sites satisfying the conditions:

\[ x_i = 2^{n-1} m_i , \quad m_i \in \mathbb{Z}, \quad i = 1, \ldots, d , \]

\[ \sum_i x_i = 2^n l + 2^{n-1} , \quad l \in \mathbb{Z} . \] (2.12)

Then the n-th step blocked fermions interact again via bilinear interactions through gauge field fat links of maximum length \(2^n\) (in units of \(a\)). One can actually do better if one allows maximum length \(2^{n+1}\) and performs a two-stage blocking of fermions interior to each blocked hypercube. The general lesson, though, is that one cannot achieve a manifestly local blocked fermion action residing on a purely hypercubic lattice, i.e. without some non-empty analog of the set \(S\) above (cf. Fig. 1). This seems to accord with previous studies of fermion actions \([11]\).

### 3. Full RG blocking

For the physical questions we are here interested in, i.e. determination of the RG flow and its fixed points, RG blockings of both fermions and gauge fields must of course be implemented. A local blocking step for fermions such as above must then be accompanied by a gauge field blocking step. It is of course possible to define a variety of gauge field blocking transformations, and several instances have been employed the literature, e.g. Swendsen, DSB or HYP-type transformations. Having defined the blocked gauge field variables, however, actually implementing the gauge blocking can generally be done only numerically. In the case of pure gauge theories, MCRG provides a tried method. Generally, given a starting action one generates a set of configurations by MC which are blocked. An effective action on the blocked lattice is then determined by measuring its coupling on the blocked ensemble. Effective actions that reproduce expectations of observables over a wide range of scales can in principle be so constructed \([12] - [13]\). This procedures may be repeated over several blocking steps. If one is not interested in the effective action flow but only in observables, the two-lattice matching method \([5]\) may be used. Here one generates two ensembles of configurations, starting from two different couplings in the original action; a matching procedure is then applied to them by measuring observables on them.

In the presence of fermions, however, it is not clear exactly how such MCRG procedures are to be applied. One cannot generate a set of configurations to be blocked without first integrating out the fermions completely. One may then apply MCRG to this ensemble. In particular the two-lattice matching method has been applied in this manner in \([4] - [7]\). Eliminating the fermions of course generates a very non-local action for the gauge fields due to the resulting fermion determinant in the gauge field background, and naive universality notions do not necessarily hold. This raises some issues. It is not immediately apparent that measurements on repeatedly blocked configurations generated from this non-local action are equivalent to measurements using the local effective action in terms of the blocked fermion and gauge variables resulting from local blocking steps. In particular, it is not clear that for fermionic observables naively written in terms of simple fermion degrees of freedom on the blocked lattice sites some kind of universality guarantees equivalence to
measurements with the local blocked action and observables in terms of blocked fermion degrees of freedom. Devising ways to test assumed equivalence of procedures, or designing new MCRG procedures for full fermion-plus-gauge field RG blocking still remain a largely unexplored area.

Here, in order to get some insight, we opt for the modest goal of devising and trying out an approximate decimation scheme that can easily be explicitly carried out.

4. Approximate RG scheme

To obtain such a scheme we proceed as follows. Eliminate first the fermions on \( S \) and compensate by adjusting the hopping parameter of the remaining fermions residing on the hypercubic lattice - this may be viewed as a typical fermion potential moving approximation to get a purely hypercubic action with ‘renormalized’ gauge interactions (Fig. 2). The renormalized hopping parameter is determined by requiring that some long-distance gauge-invariant fermion correlation function remains invariant under the fermion blocking step. Thus, taking the 2-point function, \( \bar{\psi}_n U[C_{nm}] \psi_m \), (other choices, such as a meson-meson correlator, will do equally well) this requirement translates into the exact relation:

\[
\kappa(2a) = D(\kappa(a)) \kappa(a)^2,
\]

where

\[
D(\kappa(a)) = \frac{\sum_{\mu, \nu} \langle \bar{\psi}_n (1 - \gamma_\mu)(1 - \gamma_\nu) U[C_{nm}] U_\mu(n) U_\nu(n + \hat{\mu}) \psi_m \rangle}{\sum_{\mu} \langle \bar{\psi}_n (1 - \gamma_\mu) U[C_{nm}] U_\mu(n) U_\mu(n + \hat{\mu}) \psi_m \rangle}.
\]

(4.1)-(4.2) provide a recursion relation relating the hopping parameters at lattice spacings \( a \) and \( 2a \), which may, in particular, be solved to obtain the critical hopping parameter. One may then proceed to carry out the gauge field blocking step by whatever method is chosen.

The above approximation maintains bilinear fermionic interactions at each successive blocking step. This suggests the following considerations. Assume one implements some RG blocking scheme. Denoting by \( U^{(n)}, \bar{\psi}^{(n)}, \psi^{(n)} \) the blocked variables at the \( n \)-th step, then, quire generally,
one has
\[
\int_{\Lambda^{(n)}} DU^{(n)} D\psi^{(n)} D\bar{\psi}^{(n)} \exp(-S^{(n)}_{G} - S^{(n)}_{F})
= \int_{\Lambda^{(n+1)}} DU^{(n+1)} D\psi^{(n+1)} D\bar{\psi}^{(n+1)} \exp(-S^{(n+1)}_{G} - S^{(n+1)}_{F}) .
\]

If, furthermore, as after the above approximation, \( S^{(n)}_{F} \) and \( S^{(n+1)}_{F} \) maintain the same bilinear form on \( \Lambda^{(n)} \) and \( \Lambda^{(n+1)} \), respectively, then
\[
\exp(-S^{(n+1)}_{G}) = \int_{\Lambda^{(n)}} DU^{(n)} F(U^{(n)}, U^{(n+1)}) \frac{\text{Det} S^{(n)}_{F}}{\text{Det} S^{(n+1)}_{F}} \exp(-S^{(n)}_{G}) ,
\]
with \( F(U^{(n)}, U^{(n+1)}) \) specifying the blocking relation between \( U^{(n)} \) and \( U^{(n+1)} \)'s. The ratio of determinants in (4.4) can now be expected to be reasonably local. Indeed, major part of non-locality is manifestly cancelled in the ratio. This can be seen from the formal expansion of the determinants in terms of gauge field loops:
\[
\text{Det} S^{(n)}_{F} = \prod_{C} \det (1 - \kappa^{C}[M[C]]) ,
\]
where \( M[C] \) denotes the path ordered product along the loop \( C \) of the matrices \( M(U) \) defining the fermion bilinear operator in \( S^{(n)}_{F} \), the product in (4.5) being over all loops. All loops in the expansion of \( \ln \text{Det} S^{(n+1)}_{F} \) then are contained in that of \( \text{Det} S^{(n)}_{F} \). The main contribution to the ratio comes in fact from loops inside each blocked hypercube. This suggest that one may approximate it by just such a set.

4.1 Simplest RG implementation model

These considerations lead us to an approximate implementation of RG decimations of the full system of fermions and gauge fields given by the following recipe.

After a fermion blocking step, apply the approximation depicted in Fig. 2(a) combined with (4.1)-(4.2). Then the gauge field action on the spacing \( 2a \) lattice is given by (4.4). As just explained, after the cancellation of the denominator by the numerator in the determinant ratio expressed in terms of gauge field loops, we can approximate the remainder by the contributions of loops interior to each blocked hypercube - in particular, by just the single plaquette loop weighted by a parameter \( \eta \). (One may of course consider approximations involving more interior loops, e.g. \( 2 \times 2 \) loops, and more decimation parameters.) It remains to specify and perform a gauge field blocking over each hypercube. This we do by standard potential moving decimation: every interior plaquette is moved to the hypercube boundary with the appropriate weights \( 2^{d-2} \) (Fig. 2(b)). Inserting these steps in (4.4) one obtains
\[
\exp(-S^{(n+1)}_{G}[U[p^{(n+1)}]]) = \int \prod_{b \in \text{int} p^{(n+1)}} DU_{b} \det(1 - \kappa^{4} M[p^{(n)}])^{2d} \prod_{\rho^{(n)} \in p^{(n+1)}} \left( \exp(-S^{(n)}_{G}[U[p^{(n)}]]) \right)^{2d-2}
\]
giving the plaquette action for the gauge field \( S^{(n+1)}_{G}[U[p^{(n+1)}]] \) on lattice \( \Lambda^{(n+1)} \) in terms of the theory on \( \Lambda^{(n)} \). In (4.6) \( p^{(n)} \in p^{(n+1)} \) signifies the plaquettes \( p^{(n)} \) in \( \Lambda^{(n)} \) tiling plaquette \( p^{(n+1)} \).
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|                | \(N_L\) | \(N_U\) |
|----------------|--------|--------|
| SU(2) fund     | 9      | 11     |
| SU(2) adjoint  | 2      | 3      |
| SU(3) fund     | 11 - 12| 17     |
| SU(3) sextet   | 2      | 4      |

Table 1: Conformal window lower and upper bounds as obtained from the blocking recursion relations described in the text (section 4.1).

In \(\Lambda^{(n+1)}\), and the integrations are over those bonds \(b \in \partial p^{(n)}\) that also belong to the interior of \(p^{(n+1)}\). The plaquette \(p_0^{(n)}\) is one of the tiling plaquettes (which one is immaterial by symmetry and translation invariance). Note that lattices \(\Lambda^{(n)}\) and \(\Lambda^{(n+1)}\) have lattice spacings differing by a scale factor of 2. To work out (4.6) explicitly, all factors are expanded in group characters, viz.

\[
\exp(-S_{G}^{(n)}[U[p^{(n)}]]) = \left[ c_0(n) + \sum_j c_j(n)\chi(U[p^{(n)}]) \right] \\
= \exp \left[ \beta_0(n) + \sum_j \beta_j(n)\chi(U[p^{(n)}]) \right],
\]

and similarly for \(S_{G}^{(n+1)}[U[p^{(n+1)}]]\) and the determinant factor. Inserting in (4.6), on the r.h.s. the various factor powers may be evaluated by the CG relations for combining characters and the integrations may be carried out exactly by using the character orthogonality relations. As a result one obtains explicit, albeit lengthy and ugly, recursion relations determining the expansion coefficients \(c_j(n+1)\) at the \((n+1)\)-th blocking step in terms of those at the \(n\)-th step. The procedure may then be iterated any number of times. It is important to note that the action at each step in general contains an infinite number of non-trivial characters, with corresponding couplings \(\beta_j(n)\) (cf. (4.8)). They will be generically generated just after a single blocking step even if one starts with a single representation action, e.g. the Wilson fundamental representation action, on the original lattice.

This RG recursion scheme depends only on two quantities: the parameter \(\eta\), and, at each blocking step, the critical \(\kappa(2^na)\). We have applied it to \(SU(2)\) and \(SU(3)\). The results are shown in Table 1. \(N_L\) and \(N_U\) denote the number of flavors at which an IR fixed point first appears and that at which asymptotic freedom is lost, respectively. The parameter \(\eta\) was fixed by the conformal window upper bound value \(N_U\) for \(SU(2)\) fundamental representation fermions. This decimation parameter could in principle be tuned independently in each case, but it turns out that once set by the top entry in the second column, no further adjustment was found necessary in order to produce the rest of the entries with the correct \(N_U\) values. Having fixed the parameter \(\eta\) the only uncertainties in practice are in obtaining the critical \(\kappa\) values. They were obtained from estimates from (4.2) and/or values from simulations reported in the literature \([14]\). The resulting recursion relations for the character coefficients can then be run, very cheaply, to essentially arbitrary accuracy. It is noteworthy that such an approximate decimation scheme can already produce results such as shown in Table 1.
5. Discussion - Outlook

Elucidating the phase diagram of gauge theories with different fermion content is a fully non-perturbative problem that presents many challenges. All methods that have been tried so far have their practical or conceptual shortcomings. Here we have considered what should be conceptually the most straightforward approach, i.e. the flow of the system under successive RG blocking transformations.

We have first explored locality-preserving RG blocking schemes for light fermions. We saw that blockings maintaining locality in an arbitrary gauge field background can be devised, but they generically require non-hypercubic blocked fermion actions. Such schemes may prove useful in investigating a variety of interesting issues and applications, e.g. the ‘rooting’ of fermion determinants; or improvements in the derivation of effective actions for exploring the QCD phase diagram via strong coupling and hopping expansions [15] - [16] combined with simulations.

Full (fermion plus gauge field) RG blocking would seem to provide the most direct and reliable way for exploring the IR phase structure with varying fermion content. Carrying out the gauge field blocking explicitly in the presence of the blocked fermions, however, presents new challenges. In particular, implementation of MCRG procedures, which have been applied in the pure gauge theory case, is no longer straightforward for the reasons outlined above. Much work remains to be done to sort out these issues.

At the same time judiciously chosen simplifications lead to approximate schemes which prove, as we saw, surprisingly effective. We found, in particular, that combining fermion blocking with gauge blocking within a potential moving approximation yields very reasonable results compared to other computationally much more expensive methods that have being tried. It is interesting to consider why this might be so. As it is well-known in the pure gauge case potential moving gives rather accurate beta functions at weak coupling (within 2% for SU(2)). When fermions are coupled to the system, appropriate weighing of the fermion contribution by a decimation parameter (such as the parameter $\eta$ above) apparently can preserve this feature. This ensures that the upper limits of conformal windows as predicted by perturbation theory are correctly reproduced. It generally sets the flow ‘on the right path’. It is very important in this connection that this flow moves off into an infinite dimensional subspace of couplings (cf. (4.8)) - we know from MCRG studies [12], that including enough characters in the effective action is important for good behavior over a wider range of scales. Moving in a sufficiently large, and preferably infinite dimensional, coupling space is crucial for evading lattice artifacts such as critical points and boundaries that can appear when projecting into a small finite-dimensional coupling constant subspace.

If one is interested only in locating fixed points, rather than in detailed comparisons of the behavior of observables, following the RG flow provides a ‘robust’ method in the following sense. As long as the flow manages to be within the right domain of attraction, it will lead to the corresponding fixed point; high accuracy in obtaining an effective action flow is not required. Approximate blocking schemes that incorporate the right features necessary to achieve this are then perfectly adequate for the purpose. This appears to indeed be the case for the scheme considered here. This minimal scheme, and other similarly constructed decimation schemes, are amenable to refinement by the inclusion of additional decimation parameters in more elaborate blocking definitions. It remains to be seen what improvements, if any, can be achieved by such refinements.
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