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A local moment approach to the degenerate Anderson impurity model

Martin R Galpin, Anne B Gilbert and David E Logan

Chemistry Department, Physical and Theoretical Chemistry Laboratory, Oxford University, South Parks Road, Oxford OX1 3QZ, UK

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Abstract

The local moment approach is extended to the orbitally degenerate (SU(2N)) Anderson impurity model (AIM). Single-particle dynamics are obtained over the full range of energy scales, focusing on particle–hole symmetry in the strongly-correlated regime where the onsite Coulomb interaction leads to many-body Kondo physics with entangled spin and orbital degrees of freedom. The approach captures many-body broadening of the Hubbard satellites and recovers the correct exponential vanishing of the Kondo scale for all N, and its universal scaling spectra are found to be in very good agreement with numerical renormalization group (NRG) results. In particular the high-frequency logarithmic decays of the scaling spectra, obtained here in closed form for arbitrary N, coincide essentially perfectly with available numerics from the NRG. A particular case of an anisotropic Coulomb interaction, in which the model represents a system of N 'capacitively coupled' SU(2) AIMs, is also discussed. Here the model is generally characterized by two low-energy scales, the crossover between which is seen directly in its dynamics.

1. Introduction

The Anderson impurity model (AIM) [1] plays a central role in our understanding of correlated-electron physics. It has long been used to explain the properties of a diverse range of systems where strong, local Coulomb interactions are paramount [1], most recently in the area of electronic transport through nanostructures such as semiconducting quantum dots, carbon nanotubes and single-molecule devices [2, 3].

The original AIM [4] describes a system of spin-1/2 electrons, tunnelling between a metallic conduction band and a single impurity level on which they experience a local Coulomb repulsion ‘U’. This interaction renders the problem highly non-trivial, but the physics of the model has been uncovered over several decades by a range of techniques, see e.g. [1, 5–19]. Most significantly, the Kondo effect—the quenching of the impurity spin via formation of a complex many-body state between the impurity and conduction band—is now rather well understood.

In this paper, we consider the SU(2N) generalization of the AIM [1], in which the spin-1/2 electrons of the original model are replaced by particles with 2N internal ‘flavour’ degrees of freedom. As well as being important from a theoretical point of view, the N = 2 case in particular has direct relevance to recent experiments on carbon nanotube quantum dots [20–24]. The flavour index of the particles here corresponds to a combination of electron spin and orbital indices, and the model exhibits an SU(4) Kondo effect in which spin and orbital degrees of freedom are entangled [25, 26].

The SU(2N) AIM has appeared in the literature in many guises over the years. In the U → ∞ limit, where the impurity can be at most singly occupied, the model becomes somewhat simpler to analyse. It is integrable, and hence can be studied using the Bethe ansatz [9, 27–29]. Various approximations falling under the umbrella of ‘large-N expansions’ [10–16] have also been developed. These become exact in the limit N → ∞, where their thermodynamics agree with the Bethe ansatz results [1]. Generalizing large-N techniques to finite-U is however complicated [30–35], rendering this still an active area of research today [36]. Recent progress has also been made in a different direction using a slave-rotor mean-field theory [37], which recovers the correct exponential form of the Kondo scale for arbitrary N at the particle–hole symmetric limit.

In addition to analytical approaches, SU(2N) models have also been studied numerically. Thermodynamics and dynamics have been calculated using quantum Monte Carlo (QMC), see e.g. [38]. While QMC can in principle handle any degeneracy N, interaction strength U and temperature
T \cite{38}, the method is computationally rather expensive, and cannot reach the exponentially small low-energy scales on which the Kondo effect is fully manifest. The numerical renormalization group (NRG) \cite{7,8,39} has also of course been applied with great success to the SU(2) model (for detailed reviews see e.g. \cite{1,39}), as well as to finite-U SU(4) AIMS, e.g. \cite{24,40,43}. NRG is numerically exact down to the lowest-energy scales, although the scaling with $N$ of its CPU and computer memory usage is such that at present it is realistically limited in practice to the SU(2) and SU(4) cases.

Here we analyse the SU(2N) model using the ‘local moment approach’ (LMA), considering explicitly the particle–hole symmetric limit where the impurity is occupied by $N$ particles. The LMA—developed originally for the SU(2) AIM \cite{19,44,49} and subsequently extended to encompass both the pseudogap \cite{50,51,52,53} and gapped \cite{54,55} AIMS, as well as lattice-based models within dynamical mean-field theory \cite{56,57,58,59,60,61}—has been shown to circumvent many of the traditional deficiencies of previous many-body approaches. It can handle all interaction strengths, from the non-interacting limit up to the ‘strong-coupling’ regime where electron correlations dominate; and recovers the correct physics at the lowest-energy scales. It is also physically transparent and computationally straightforward; many of its results can in fact be obtained analytically, including non-trivial closed formulae for single-particle dynamics of the model in the strong-coupling regime.

We explain in the following that generalization of the LMA to the SU(2N) AIM is quite natural (and does not require significant additional computation when applied to arbitrary $N$). In the strong-coupling regime, $N$ essentially appears only as an additional prefactor to the LMA self-energies, and hence many of the known LMA results for the SU(2) model can be extended quite straightforwardly. In particular, we show that the LMA recovers the correct exponential dependence of the Kondo scale on $N$, and derive a closed expression for the ‘tails’ of the universal SU(2N) single-particle scaling spectrum that agrees excellently with NRG results for the SU(2) and SU(4) cases.

The paper is structured as follows. Section 2 defines the SU(2N) AIM, shows its connection to a model of $N$ capacitive coupled SU(2) AIMS, and defines the particle–hole symmetric case on which we focus here. In section 3 we introduce the LMA, starting with an examination of the non-interacting and ‘atomic’ limits, before moving on to the mean-field level of unrestricted Hartree–Fock and then the LMA itself. Section 4 contains a summary of our results for the model, both numerical and analytic as mentioned above, together with direct comparison to NRG results for single-particle dynamics in the SU(4) and SU(2) cases. We end the paper by discussing briefly the physics of the model away from SU(2N) symmetry (section 5), and provide a short conclusion in section 6.

2. Model

In conventional notation the Hamiltonian for the SU(2N) Anderson impurity model is \cite{1}

$$\hat{H} = \sum_{k,m} \epsilon_{k} c_{k\alpha m}^{\dagger} c_{k m}^{\dagger} + \sum_{i} n_{i\sigma}^{\dagger} + U \frac{1}{2} \sum_{m,n} n_{m\sigma}^{\dagger} n_{n\sigma}^{\dagger} + V \sum_{k,m} (c_{k m}^{\dagger} f_{m} + \text{h.c.}), \quad (1)$$

where $n_{m\sigma} = f_{m\sigma}^{\dagger} f_{m\sigma}$, the flavour index $m$ takes on $2N$ distinct values, and the prime in the third sum means $m' = m$ is excluded. The Hamiltonian thus describes a non-interacting host band of ‘flavourful’ particles (first term), plus a localized impurity orbital with onsite Coulomb repulsion $U$ (second and third terms), with tunnel couplings between the two (final term).

Equation (1) highlights clearly the inherent SU(2N) symmetry of the model. But it will also be instructive to rewrite the Hamiltonian in a way that reveals its connection to a problem of spinful SU(2) electrons. To this end, we regard $m$ as a composite index, $m \equiv (i, \sigma)$, where $i$ takes on $N$ integral ‘site’ indices from 1 to $N$ and $\sigma = \uparrow$ or $\downarrow$ represents an SU(2) spin index. Sums over $m$ then become joint sums over $i$ and $\sigma$, and (1) can be rewritten as

$$\hat{H} = \sum_{k,i,\sigma} \epsilon_{k i\sigma} c_{k i\sigma}^{\dagger} c_{k i\sigma}^{\dagger} + \sum_{i,\sigma} \tilde{n}_{i\sigma} + \frac{U}{2} \sum_{i,j,\sigma,\sigma'} \tilde{n}_{i\sigma}^{\dagger} \tilde{n}_{j\sigma'}^{\dagger} + V \sum_{k,i,\sigma} (c_{k i\sigma}^{\dagger} f_{i\sigma} + \text{h.c.}) \quad (2)$$

with $\tilde{n}_{i\sigma} = f_{i\sigma}^{\dagger} f_{i\sigma}$, and the prime on the sum means that the term with both $i = j$ and $\sigma = \sigma'$ is excluded. The interaction term can be simplified by writing

$$\sum_{i,j,\sigma,\sigma'} \tilde{n}_{i\sigma}^{\dagger} \tilde{n}_{j\sigma'}^{\dagger} \equiv \sum_{i,j} \delta_{ij} + \sum_{i,j} \delta_{\sigma,\sigma'} \quad (3)$$

(with $\delta_{ab}$ the Kronecker delta), such that

$$\hat{H} = \sum_{i} \left[ \sum_{k,m} \epsilon_{k} c_{k i\sigma}^{\dagger} c_{k i\sigma}^{\dagger} + \epsilon_{i} \tilde{n}_{i\sigma} + U \tilde{n}_{i\sigma}^{\dagger} \tilde{n}_{\sigma}^{\dagger} \right] + V \sum_{k,\sigma} (c_{k i\sigma}^{\dagger} f_{i\sigma} + \text{h.c.}) \quad \left[ \sum_{i,j} \tilde{n}_{i\sigma}^{\dagger} \tilde{n}_{j\sigma}^{\dagger} \right] + \frac{U}{2} \sum_{i,j} \tilde{n}_{i\sigma}^{\dagger} \tilde{n}_{j\sigma}^{\dagger} \quad (4)$$

where $\tilde{n}_{i\sigma} = \sum_{\sigma} n_{i\sigma}$. In this form, the Hamiltonian describes $N$ equivalent capacitively coupled SU(2) Anderson models, in the limit of identical onsite and intersite Coulomb interactions. The LMA developed here can in fact be straightforwardly generalized to the case when the intersite interaction is less than $U$; we comment briefly on the main results of such in section 5, but until then focus on the fully symmetric SU(2N) limit corresponding to (1).

For relative simplicity we also consider the model in the particle–hole symmetric limit, with a standard flat, infinitely wide conduction band with density of states $\rho_{0} \left[1\right]$. The latter automatically ensures particle–hole symmetry of the host; the impurity part of (1) under a particle–hole transformation $f_{m} \rightarrow$
the approach below to the particle–hole asymmetric SU3.1. Non-interacting limit as particular cases. The physics of the full model are seen in these rather trivial limits, ϵform, in direct analogy to the Δ1 limit with Vm to the particle–hole symmetric limit adds here that the subsequent approach is not in fact restricted away from particle–hole symmetry [44], and generalization of this way. For later use, note that the Hamiltonian (1) in this limit reduces (modulo an irrelevant constant) to the capacitive charging form

\[ \hat{H} = \sum_{k,m} \epsilon_k c_{km}^+ c_{km} + \frac{U}{2} \sum_{m,m'} \hat{n}_m \hat{n}_{m'} \] (7)

with \( \hat{N} = \sum_m \hat{n}_m \) the total impurity charge operator. We add here that the subsequent approach is not in fact restricted to the particle–hole symmetric limit per se; the LMA has already been used successfully to describe the SU(2) AIM away from particle–hole symmetry [44], and generalization of the approach below to the particle–hole asymmetric SU(2N) model can be developed in a similar vein.

To examine the dynamics of the SU(2N) AIM, the central quantity of interest is the time-ordered, flavour-m impurity Green function \( G_m(\omega) \), defined as usual [62] by the Fourier transform of \( G_m(t) = -i \langle \hat{\cal T} \{ f_m(t) f_m^0(0) \} \rangle \) (with \( \hat{\cal T} \) the Wick time-ordering operator), and with corresponding single-particle spectrum \( D_m(\omega) = -\pi^{-1} \text{sgn}(\omega) \text{Im} G_m(\omega) \). It is this on which we naturally focus in the paper.

3. Local moment approach

Before we develop the LMA for the SU(2N) AIM, it is instructive to comment briefly on two special cases: the non-interacting limit of \( U = 0 \) (but arbitrary \( V \)), and the atomic limit with \( V = 0 \) (but arbitrary \( U \)). While only glimpses of the physics of the full model are seen in these rather trivial limits, our main purpose in considering them is that any credible approximate theory for the model should recover both limits as particular cases.

3.1. Non-interacting limit

In the non-interacting limit, \( U = 0 \), the impurity Green function (here denoted by \( g_m(\omega) \)) is readily obtained in closed form, in direct analogy to the SU(2) case [1], namely

\[ g_m(\omega) = \frac{1}{\omega^+ - \Delta(\omega)} \] (8)

with \( \omega^+ = \omega + i0^+ \text{sgn}(\omega) \). Here \( \Delta(\omega) = \sum_k |V_k|^2/(\omega^+ - \epsilon_k) \) is the host-impurity hybridization function, reducing to \( \Delta(\omega) = -i\text{sgn}(\omega) \Delta_0 \) for the conventional wide-band host considered explicitly, where \( \Delta_0 = \pi \rho_0 |V|^2 \) (with \( \rho_0 \) the conduction band density of states, as above). The spectrum of (8) is simply

\[ \pi \Delta_0 d_m(\omega) = \frac{\Delta_0^2}{\omega^2 + \Delta_0^2}; \] (9)

the Lorentzian form of which reflects the lifetime broadening (of width \( \Delta_0 \)) of the impurity level due to its hybridization with the host band. This introduces a natural energy scale to the problem, and it is convenient to use \( \Delta_0 \) as the unit of energy in the rest of the paper (unless stated otherwise). To this end, we define the reduced Coulomb interaction \( \tilde{U} = U/\Delta_0 \) and the reduced frequency \( \tilde{\omega} = \omega/\Delta_0 \).

3.2. Atomic limit

The atomic limit (\( V = 0 \)) is trivially soluble for any value of \( U \), since the impurity number operator \( \hat{N} \) commutes with \( \hat{H} \). The basic physics follows directly from the Hamiltonian in (7).

When \( U > 0 \), the ground state is necessarily degenerate. The impurity is occupied by \( N \) particles in its ground state, and these can be distributed amongst the 2N orbitals in \( W_N = (2^N) \) distinct ways. Single-particle excitations connect the degenerate ground state manifold to further \( N \pm 1 \)-particle degenerate manifolds, all lying \( U/2 \) higher in energy.

In the absence of any special preparation, the particular ground state adopted by the system is equally likely to be any one of the \( W_N \) possibilities. The single-particle excitation spectrum for a given flavour \( m \) is then the statistical average of the individual flavour-m spectra of all the \( W_N \) ground states. In precisely \( W_N/2 \) of these, there will be a particle in flavour orbital \( m \) and hence a ‘removal’ pole in the spectrum at \( \omega = -U/2 \). In the other \( W_N/2 \) cases the \( m \)th flavour orbital will be unoccupied, leading to an ‘addition’ pole at \( \omega = +U/2 \). As a result, the spectrum in the atomic limit consists of two poles of weight \( 1/2 \) lying at \( \omega = \pm U/2 \), i.e.

\[ G_m(\omega) = \frac{1}{2} \left[ \frac{1}{\omega^+ + \frac{U}{2}} + \frac{1}{\omega^+ - \frac{U}{2}} \right], \] (10)

independently of \( N \).

3.3. Mean field

Next we turn to an analysis at static mean-field (MF) level. By analogy to unrestricted Hartee–Fock (UHF) for the SU(2) case [4], we allow for the most general form of local symmetry-breaking, writing \( \hat{n}_m = \langle \hat{n}_m \rangle + \delta \hat{n}_m \) for all flavour indices \( m \). Substituting this into (1) and neglecting the fluctuation terms quadratic in \( \delta \hat{n}_m \delta \hat{n}_{m'} \) gives, to within an irrelevant constant,

\[ \hat{H} = \sum_{k,m} \epsilon_k c_{km}^+ c_{km} + \sum_m \epsilon_m \hat{n}_m + V \sum_{k,m} c_{km}^+ f_m + \text{h.c.}. \] (11)

with

\[ \epsilon_m = \epsilon + U \sum_{m' \neq m} \langle \hat{n}_{m'} \rangle. \] (12)

Under a particle–hole transformation \( \hat{\cal N} \rightarrow 2N - \hat{\cal N} \), and since \( \hat{H} \) is invariant under this transformation (see (7)), particle–hole symmetry implies the single constraint

\[ \sum_m \langle \hat{n}_m \rangle = N, \] (13)

\(^1\) All energies are measured with respect to the Fermi level.
so that there are $2N - 1$ possible channels in which symmetry can be broken at MF level. Use of (13) in (12), along with (6), yields an expression for $\epsilon_m$ in terms of $\langle \hat{n}_m \rangle$ alone, namely

$$\epsilon_m = U\left( \frac{1}{2} - \langle \hat{n}_m \rangle \right).$$  (14)

Since the Hamiltonian in (11) is separable for each flavour $m$, the MF impurity Green functions follow in direct analogy to (8) as

$$G_m(\omega) = \frac{1}{\omega + \epsilon_m + i \text{sgn}(\omega) \Delta_0}.$$  (15)

The mean number of flavour-$m$ particles on the impurity is then given by integrating the spectrum $D_m(\omega) = -\pi^{-1} \text{sgn}(\omega) \text{Im} G_m(\omega)$ up to the Fermi level:

$$\langle \hat{n}_m \rangle = \int_{-\infty}^{0} D_m(\omega) d\omega = \frac{1}{2} - \frac{1}{\pi} \tan^{-1}\left( \frac{\epsilon_m}{\Delta_0} \right).$$  (16)

Together with (14), this implies the following MF self-consistency equation for each $\langle \hat{n}_m \rangle$:

$$\langle \hat{n}_m \rangle = \frac{1}{2} - \frac{1}{\pi} \tan^{-1}\left( \frac{U\left( \frac{1}{2} - \langle \hat{n}_m \rangle \right)}{\Delta_0} \right).$$  (17)

subject to the constraint of (13).

The simplest solution of (17) and (13) arises when $\langle \hat{n}_m \rangle = \frac{1}{2}$ for all $m$. This is restricted Hartree–Fock (RHF), in which the SU(2N) symmetry of the impurity is left entirely unbroken. In this case, by (12), all the $\epsilon_m$s vanish, and hence (15) reduces to the non-interacting Green function $g_m(\omega)$ (8) for all $m$. But while RHF is a (trivial) self-consistent solution of the MF equations, it is not the solution of lowest energy. By breaking the symmetry of (1), variationally lower solutions can be generated [4].

We have analysed the broken-symmetry MF solutions in detail. As now shown, the self-consistent solutions have the $2N$ mean-field energies $\epsilon_m$ partitioned into two groups of $N$ degenerate levels. We label the energy levels of the two groups as $\epsilon_+$ and $\epsilon_-$, and the mean occupation of each level as $\langle \hat{n}_+ \rangle$ and $\langle \hat{n}_- \rangle$, respectively. For convenience in what follows, we simply refer to the two groups of $N$ levels as the ‘+’ and ‘−’ groups, respectively.

Upon partitioning the levels between these two groups, the $2N$ equations in (17) reduce to just two:

$$\langle \hat{n}_\pm \rangle = \frac{1}{2} - \frac{1}{\pi} \tan^{-1}\left( \frac{U\left( \frac{1}{2} - \langle \hat{n}_\pm \rangle \right)}{\Delta_0} \right),$$  (18)

and the constraint in (13) reads $\sum_m \langle \hat{n}_m \rangle = N\langle \hat{n}_+ \rangle + N\langle \hat{n}_- \rangle = N$, i.e.

$$\langle \hat{n}_+ \rangle + \langle \hat{n}_- \rangle = 1.$$  (19)

This pair of self-consistency equations (18) and (19), is formally identical to that of the SU(2) AIM at UHF level [4]. In addition to the RHF solution described above ($\langle \hat{n}_\pm \rangle = \frac{1}{2}$), a symmetry-broken solution is thus found to arise [4] whenever $\hat{U} > \pi$: physically the ‘+’ and ‘−’ groups are split about $\omega = 0$, leading to a net lowering of the energy. The splitting of the levels is described most easily by introducing the ‘moment’ parameter $\mu$;

writing $\langle \hat{n}_\pm \rangle = \frac{1}{2}(1 \mp \mu)$ (which automatically satisfies (19)). The remaining self-consistency equation (18), then reads

$$\mu = \frac{2}{\pi} \tan^{-1}\left( \frac{\hat{U}\mu}{2} \right),$$  (20)

from which it follows that symmetry-broken solutions with $\mu \neq 0$ arise in degenerate pairs; namely $\mu = \pm |\mu_0|$ and $-|\mu_0|$, which we denote respectively as the ‘A’-type and ‘B’-type solutions. From (14) one sees that an ‘A’-type solution has the energy levels at $\epsilon_{\pm} = \pm \frac{1}{2}U|\mu_0|$, while the ‘B’-type solution by contrast has $\epsilon_\pm = \mp \frac{1}{2}U|\mu_0|$.

For a given type of solution (A or B), the mean-field ground state is degenerate. The $2N$ mean-field impurity levels can be partitioned between two groups of $N$ levels in $W_N = \binom{N}{N}$ distinct ways. For example, one solution corresponds to allocating flavours $m = 1–N$ to the ‘−’ group and the remaining flavours $m = (N + 1)–2N$ to the ‘+’ group; a second solution arises when the two sets of flavour components are completely interchanged; and other solutions can obviously be obtained by interchanging only some of the flavour components between the two groups. Bearing this in mind, one can if desired enumerate all $W_N$ possible degenerate MF states for any given $N$, labelling them using the following notation. For each state, the $2N$ possible flavour components are written in a row, such that the first $N$ belong to the ‘−’ group and the remaining $N$ are those of the ‘+’ group. For example, in the case $N = 2$ [SU(4)], the state (1 2 3 4) has $\epsilon_1 = \epsilon_- = \epsilon_2 = \epsilon_+ = \epsilon_4$. All other states can be generated by permutations of the flavour components: e.g. the permutation (1 3 2 4) gives the state in which $\epsilon_1 = \epsilon_- = \epsilon_3$, and $\epsilon_2 = \epsilon_+ = \epsilon_4$. Note that some permutations lead to physically equivalent states—e.g. (1 3 2 4) and (3 1 2 4)—since the ordering of levels within a group is irrelevant. To avoid this, one considers only permutations in which the two sets of $N$ integers (corresponding to the ‘−’ and ‘+’ groups) are each in ascending numerical order; i.e. the state (1 3 2 4) is allowed, whereas (3 1 2 4) is not. In this way, every distinct MF state is counted once, and once only.

The net MF Green function for flavour $m$, denoted here by $G_m^0(\omega)$, then follows by averaging the broken-symmetry, flavour-$m$ Green functions of the $W_N$ degenerate ground states:

$$G_m^0(\omega) = \frac{1}{W_N} \sum_P G_m^P(\omega).$$  (21)

where $G_m^P(\omega)$ is the flavour-$m$ Green function for MF solution $\alpha (= \text{either A or B})$ and a particular permutation $P$ of the flavour components between the $\epsilon_\pm$ and $\epsilon_\mp$ levels. The sum over $P$ is taken over the set of permutations that enumerates all $W_N$ physically distinct states, as described above. Equation (21) can then be simplified by the following arguments. First, as seen explicitly in (15), the MF $G_m(\omega)$ depends on $\epsilon_m$ only. Therefore, on defining

$$G_m^{\alpha}(\omega) = \frac{1}{\omega + \epsilon_\pm + i \text{sgn}(\omega)\Delta_0},$$  (22)

we obtain $G_m^{P}(\omega) = G_m^{\alpha}(\omega)$ if the permutation $P$ allocates flavour component $m$ to the ‘±’ group, respectively. And by
symmetry, when the \( W_N \) degenerate states are enumerated the flavour \( m \) will appear \( W_N/2 \) times in each of the ‘+’ and ‘−’ groups (\( W_N \) is necessarily even for \( N \geq 1 \)). Combining these results, one obtains

\[
G^0_\alpha(\omega) = \frac{1}{2}[G^A_\alpha(\omega) + G^B_\alpha(\omega)]
\]

(23)

for either \( \alpha = A \) or \( B \) (and obviously being independent of \( m \), reflecting \( SU(4) \) rotational invariance).

Hence, the net MF Green function is equivalently the average of the MF Green functions for the + and − groups, regardless of how the flavour components are actually allocated to those groups to form a particular MF solution. Moreover, since interconverting \( A \) and \( B \) exchanges the energies of the two groups such that \( G^\pm_\alpha(\omega) = G^\mp_\alpha(\omega) \), (23) becomes

\[
G^0_m(\omega) = \frac{1}{2}[G^A_\alpha(\omega) + G^B_\alpha(\omega)];
\]

(24)

the statistically averaged MF Green function is thus equivalently the average of the ‘A’-and ‘B’-type Green functions for either given group (+ or −), again regardless of how the flavour components are partitioned between the groups.

In either case, on taking the imaginary part and using (22), the average UHF spectrum follows as

\[
\pi \Delta_0 D^0_m(\omega) = \frac{1}{2} \left[ \left( \frac{\Delta_0^+}{(\omega - \frac{1}{2}U|\mu_0|)^2 + \Delta_0^+} \right) + \left( \frac{\Delta_0^-}{(\omega + \frac{1}{2}U|\mu_0|)^2 + \Delta_0^-} \right) \right],
\]

(25)

consisting therefore of two superimposed Lorentzians (‘Hubbard satellites’) of half-width, half-maximum (HWHM) \( \Delta_0 \).

Note that when \( \Delta_0 \) is switched off, the MF spectrum (25) reduces correctly to that of the atomic limit, itself obtained from the imaginary part of (10). This is seen directly by taking the limit \( \Delta_0 \to 0 \) in (25): the self-consistency condition of (20) becomes \( \mu = \text{sgn}(\mu) \) (and hence \( |\mu_0| = 1 \)), while the Lorentzians become delta-functions at \( \omega = \pm \frac{1}{2}U|\mu_0| \). And when \( U = 0 \), the non-interacting spectrum, (9), is trivially recovered by (25).

The UHF approach thus captures both the atomic and non-interacting limits (sections 3.1 and 3.2), and in some sense bridges the two when \( U \) and \( \Delta_0 \) are both finite. But it does not of course capture the full physics of the model. Equation (25) is independent of \( N \); in reality, the widths of the Hubbard satellites increase with \( N \), since this opens up more relaxation channels which lead to enhanced lifetime broadening [10, 19] (see also section 4.1 below). Most importantly, the low-energy description provided by (25) is entirely wrong. The spectrum of the AIM is well known to possess an exponentially narrow Kondo resonance straddling the Fermi level [1]. This many-body resonance is simply absent at the static MF level of description, and can only be captured by developing a more sophisticated approach.

In section 3.4 we describe how the local moment approach can be extended to the \( SU(2N) \) AIM. As explained previously in a number of works (see e.g. [19, 44]), the technique overcomes the intrinsic limitations of static MF theory via inclusion, within an inherent two-self-energy framework, of dynamical self-energy contributions which in physical terms embody tunnelling between degenerate MF states; and which, in acting thereby to restore the symmetry broken at pure MF level, lead correctly to recovery of Fermi-liquid behaviour on low-energy scales.

To set up the LMA in practice, it is convenient to work with the ‘capacitively coupled’ quantum dot picture of (4), replacing all \( m \) indices by \( (i, \sigma) \). And just as the MF Green function \( G^0_{\alpha}(\omega) \) ((23) or (24)) is independent of the MF solution with which one chooses to work, the same can readily be shown to hold within the LMA. We thus choose to work with the particular MF solution in which all \( \sigma = \uparrow \) levels are allocated to the ‘−’ group, and all \( \sigma = \downarrow \) levels are allocated to the ‘+’ group. Physically, this solution corresponds to uniform charges on all sites \( i \), but broken spin-symmetry; more precisely, when \( |\mu| > 0 \) the ‘A’-type solution has an excess of \( \uparrow \)-spin electrons on every (equivalent) site, while the ‘B’-type solution has an excess of \( \downarrow \)-spins. Other partitionings of the flavour components \( (i, \sigma) \) would of course correspond to different physical pictures—such as MF solutions that are spin-symmetric but of broken charge symmetry—but we reiterate that which is used in practice is irrelevant when calculating the full rotationally invariant Green function. Writing (23) and (24) in the chosen charge-symmetric, broken spin-symmetry picture, gives

\[
G^0_{\alpha}(\omega) = \frac{1}{2}[G^0_{\alpha}(\omega) + \tilde{G}^0_{\alpha}(\omega)]
\]

(26)

\[
= \frac{1}{2}[G^0_{\alpha}(\omega) + G^0_{\beta}(\omega)]
\]

(27)

(where \( \alpha = \text{either A or B} \), which will be employed in due course; with symmetries \( G^0_{\alpha}(\omega) = G^0_{\beta}(\omega) \), such that from (27) \( G^0_{\alpha}(\omega) \) is independent of spin \( \sigma \) (and as such spin-rotationally invariant [19, 49]), as well as being independent of \( i \) (reflecting the equivalence of the \( N \) levels in the capacitively coupled quantum dot picture).

3.4. Local moment approach

The MF solutions above provide a natural starting point for perturbation theory within a two-self-energy description [19, 44, 49]. We take the unperturbed Hamiltonian to have the same symmetries as that of MF, i.e. to be of form

\[
\hat{H}^0 = \sum_{k,i,\sigma} \epsilon_k c_{k\sigma}^\dagger c_{ki\sigma} - \sum_{i,\sigma} \alpha_\sigma x \hat{n}_{i\sigma} + V \sum_{k,i,\sigma} (c_{ki\sigma}^\dagger f_{i\sigma} + \text{h.c.})
\]

(28)

for each of \( \alpha = \text{A or B} \), taking \( A \equiv + \) and \( B \equiv - \) for notational convenience when using the label \( \alpha \); and where \( x = U|\mu|/2 \), with local moment \( |\mu| \) (determined at post-MF level via the symmetry restoration condition intrinsic to the LMA [44], and specified at the end of the section). The perturbation term corresponding to (28) is \( \hat{H}^1 = \hat{H} - \hat{H}^0 \), namely

\[
\hat{H}^1 = \sum_i U \hat{n}_{i\uparrow} \hat{n}_{i\downarrow} + \frac{U}{2} \sum_{i,j} \hat{n}_{i\uparrow} \hat{n}_{j\downarrow} + \sum_{i,\sigma} (\epsilon + \alpha_\sigma x) \hat{n}_{i\sigma}.
\]

(29)

and in precise parallel to the MF results of section 3.3, the unperturbed Green functions for a \( \sigma \)-spin electron on orbital \( i \) are

\[
G^0_{\alpha}(\omega) = \frac{1}{\omega + \alpha_\sigma x + i\text{sgn}(\omega)\Delta_0}.
\]

(30)
The corresponding perturbed Green functions can be obtained by diagrammatic perturbation theory in $\hat{H}^\sigma_i$ [49]. One naturally obtains the Dyson equation

$$G^\sigma_{i\sigma}(\omega) = \frac{1}{[G^\sigma_{i\sigma}(\omega)]^{-1} - (\epsilon + \alpha \sigma \chi) - \tilde{\Sigma}_{i\sigma}^\sigma(\omega)}$$

(31)

which defines the two self-energies $\tilde{\Sigma}_{i\sigma}^\sigma(\omega)$ (for $\sigma = \uparrow, \downarrow$) arising for a given $\alpha$ (our definition of $\Sigma_{i\sigma}^\sigma(\omega)$ excluding the static first-order diagram $(\epsilon + \alpha \sigma \chi)$ coming from the third term in (29)). The full, rotationally invariant impurity Green function then follows using directly analogous arguments to those of the previous section, such that (cf (26))

$$G_m(\omega) = \frac{1}{2} \sum_{\sigma} G^\sigma_{i\sigma}(\omega)$$

(32)

(again independently of $m = (i, \sigma)$). The spin-symmetry of the Hamiltonian (28) implies that

$$G^\sigma_{i\sigma}(\omega) = G^{\bar{\sigma}}_{i\bar{\sigma}}(\omega) ;$$

while particle-hole symmetry implies

$$G_{i\alpha}(\omega) = -G_{i\bar{\alpha}}(-\omega),$$

thus simplifying the calculation of (32) in practice.

Following [19, 44, 49], we now expand the $\tilde{\Sigma}_{i\sigma}^\sigma$s perturbatively in their respective $H^\sigma_i$s. For specificity we consider $\alpha = A$ only (and for clarity drop the $A$ superscripts from here on unless indicated otherwise explicitly); the B-type self-energy follows (if desired) from the symmetry (33). As in previous LMA papers (e.g. [19, 44]), we choose to separate the self-energies into static ($\omega$-independent) plus dynamical terms. The former is approximated by the sum over ‘tadpole’ diagrams in figure 1(a), which gives

$$\tilde{\Sigma}_{i\sigma}(\omega) = \sum_{j,\sigma'} \frac{1}{2} U(1 + \sigma'|\bar{\mu}|) + \Sigma_{i\sigma}(\omega)$$

$$(N - \frac{1}{2}) U - \frac{1}{2} \sigma U |\bar{\mu}| + \Sigma_{i\sigma}(\omega)$$

(35)

(36)

and $\bar{\mu}$ is defined by

$$|\bar{\mu}| = \int_{-\infty}^{0} d\omega \ [D_{i\uparrow}(\omega) - D_{i\downarrow}(\omega)] = \frac{2}{\pi} \tan^{-1} \frac{x}{\Delta_0}$$

(37)

Generalizing the approach of [19], we approximate the dynamical part of the self-energy for a flavour-$m \equiv (i, \sigma)$ particle by the infinite series shown in figure 1(b). As we are working from the broken-symmetry states in which the $\tilde{G}_{i\sigma}$s are independent of orbital index $j$, all $N$ diagrams describing interactions with $-\sigma$-spin electrons are equivalent, as are all remaining $N - 1$ diagrams describing interactions with $\sigma$-spin electrons. The sum over $j,\sigma'$ in figure 1(b) can thus be replaced by just two terms. Upon recasting the diagrams in terms of random-phase approximation (RPA) polarization propagators, we obtain the result shown in figure 2.

Note that all explicit $N$ dependence to the self-energies is contained in the coefficients of the two diagrams, the broken-symmetry Green functions from which the diagrams are constructed being independent of explicit factors of $N$. In the $SU(2)$ case ($N = 1$), only the first diagram on the
right-hand side of figure 2(a) survives and the self-energy approximation reduces to that used in [19]. When \( N > 1 \), both diagrams contribute, although we find that the dominant contribution in the strongly-correlated regime, \( \tilde{U} \gg 1 \), comes from the first diagram alone (see appendix B).

Translating the diagrams in figure 2 using the Feynman rules gives (cf [19])

\[
\Sigma_{i\sigma}(\omega) = N \Sigma_{i\sigma}^{\sigma-\sigma}(\omega) + (N - 1) \Sigma_{i\sigma}^{\sigma \sigma}(\omega)
\]  

(38)

where

\[
\Sigma_{i\sigma}^{\sigma \sigma'}(\omega) = \frac{U^2}{2\pi i} \int_{-\infty}^{\infty} d\omega_1 \Pi_{i\sigma}(\omega_1) G_{i\sigma'}(\omega_1 + \omega). 
\]  

(39)

The RPA polarization propagators \( \Pi_{i\sigma}(\omega) \) appearing above are translated as

\[
\Pi_{i\sigma}(\omega) = \frac{\Pi_{i\sigma}^0(\omega)}{1 - U \Pi_{i\sigma}^0(\omega)}, 
\]  

(40)

with the ‘bare’ polarization bubble

\[
\Pi_{i\sigma}^0(\omega) = -\frac{1}{2\pi i} \int_{-\infty}^{\infty} d\omega_1 \ G_{i\sigma'}(\omega_1) G_{i\sigma}(\omega_1 - \omega). 
\]  

(41)

For the flat-band AIM of interest, the latter can be obtained in closed form. The equations for \( \Pi_{i\sigma}^0(\omega) \) are provided in the appendix of [19], and those for \( \Pi_{i\sigma}(\omega) \) are given in appendix A of the present work.

We note that the numerical calculation of the LMA dynamical self-energies is quite straightforward. In practice it is convenient to determine separately the real and imaginary parts of the various constituents listed above, the former obtained by Hilbert transformation of the latter. As the methods used are essentially identical to those of previous work [19], we do not discuss them further here.

It remains finally to specify the condition for determination of the local moment \( |\mu| \) (entering parametrically via \( x = U|\mu|/2 \) in the Hamiltonian (28)). This is achieved by ‘symmetry restoration’, the central element of the LMA, corresponding physically to restoration of the symmetry which is broken at pure MF level. It amounts to a single self-consistency condition on the two self-energies precisely at the Fermi level, \( \omega = 0 \); which, regardless of the specific diagrammatic approximation used for the self-energies, ensures that the rotationally invariant LMA Green function \( G_{i\sigma}(\omega) \) recovers Fermi-liquid behaviour on the lowest-energy scales [44, 49]. Since symmetry restoration is discussed extensively elsewhere (see e.g. [44, 45, 49, 52, 58]) we do not elaborate further on it here; but simply note that in direct correspondence with earlier work, the symmetry restoration condition here is

\[
\tilde{\Sigma}_{i\uparrow}(\omega = 0) = \tilde{\Sigma}_{i\downarrow}(\omega = 0).
\]  

(42)

In addition, analyticity of the RPA transverse spin polarization propagator requires [19]

\[
|\mu_0| \equiv |\mu| = \frac{2\pi U}{U} < 1. 
\]  

(43)

Equations (42) and (43) are readily solved numerically for any choice of the bare model parameters, and are sufficient to determine \( x \) uniquely in all cases.

For later use, we also note that a combination of particle–hole symmetry (34) and \( \uparrow / \downarrow \)-spin symmetry (33) implies that (42) reduces to

\[
\epsilon + \tilde{\Sigma}_{i\sigma}(\omega = 0) = 0. 
\]  

(44)

Hence, from (36) and (6), the symmetry restoration condition of (42) can be written alternatively as

\[
\Sigma_{i\sigma}(\omega = 0) = \frac{\sigma}{2} U|\tilde{\mu}|. 
\]  

(45)

4. Results

We have analysed in detail the \( SU(2N) \) AIM within the LMA described above. Numerical evaluation of the impurity Green function is computationally very inexpensive, and we have examined its behaviour over a wide range of the bare model parameters. Here we describe our main results, focusing in particular on the strong-coupling limit, \( \tilde{U} = U/\Delta_0 \gg 1 \), where the model enters the Kondo regime in which electron correlations are key.

4.1. Single-particle spectrum: overview

Figure 3(a) shows representative single-particle spectra of the \( SU(4) \) AIM in strong coupling, calculated within the LMA as described in the previous section, for \( \tilde{U} = 20, 40 \) and 50. The spectra are plotted as \( \pi \Delta_0 D(\omega) \) versus \( \omega/\Delta_0 \) on a linear frequency scale, and at the Fermi level in particular are seen to satisfy correctly the dictates of the Friedel sum rule for all \( SU(2N) \) at half-filling, \( \pi \Delta_0 D(\omega = 0) = 1 \) [1, 24]. The figure also shows for comparison the corresponding spectrum of the \( SU(2) \) AIM [19] for \( \tilde{U} = 20 \).

Consider first the high-frequency behaviour seen in figure 3(a). The spectra all show clear Hubbard satellites arising on the scale \( |\omega| \approx U/2 \), reflecting physically the single-particle excitations of the atomic limit (see section 3.2). The finite coupling to the host broadens the Hubbard satellites from the atomic-limit delta-functions in (10). This broadening is \( O(\Delta_0) \) in both cases, but the Hubbard satellites of the \( SU(4) \) spectra are significantly broader than those of the \( SU(2) \) spectrum (moreover, both are broader than the UHF spectrum in (25)). The origin of this effect, as alluded to at the end of section 3.3, is many-body scattering from the impurity site [19], which is not of course captured at pure MF level. The energy cost to excite particles between different flavour states on the impurity site (i.e. ‘flip a spin’ in the \( SU(2) \) case) is of order the Kondo scale. This is negligible on the energy scale of the Hubbard satellites, and hence the number of relaxation channels contributing to the lifetime broadening of the Hubbard satellites increases with \( N \). Indeed in the particle–hole symmetric limit considered here, it is straightforward to work out the LMA that the widths of the Hubbard satellites go as \( (N + 1)\Delta_0 \) in the strong-coupling limit, \( \tilde{U} \gg 1 \) [10, 38].

Moving to lower-energy scales, we see that the LMA indeed captures the \( SU(4) \) Kondo physics, all spectra in
While its effect on the spectrum is slight, it can be removed if so desired by the procedure described in [45].

Figure 3(a) displays a sharp Kondo resonance centred on the Fermi level $\omega = 0$. This behaviour is clearer on the much reduced linear frequency scale shown in figure 3(c); but is seen most clearly by showing the spectra on a logarithmic frequency scale as shown in figure 3(b). The evident conclusion to be drawn from figure 3(b) is that the low-energy behaviour of the three $SU(4)$ spectra (solid lines) is universal [38, 42]. Defining a characteristic Kondo scale $\omega_K$ by the HWHM of the Kondo resonance, it is quite clear that the three solid lines will scale onto each other when plotted on the reduced frequency scale $\omega/\omega_K$ (provided naturally that $\omega$ itself does not approach non-universal scales of order $|\omega| \sim O(\Delta_0)$); as considered in detail in section 4.3 below. Note further that the $SU(2)$ spectrum shown as a dotted line in figure 3(b), itself known [19, 45] to display universal scaling as a function of $\omega/\omega_K$, does not scale onto the universal $SU(4)$ behaviour: i.e. the $SU(2)$ and $SU(4)$ scaling spectra are distinct [42] (cf [38]). To understand these

To verify (46) itself, we have calculated $\omega_m$ numerically over a wide range of $\tilde{U} \gg 1$, for the $SU(2)$, $SU(4)$ and $SU(6)$ cases. The results are shown in figure 4, plotted as $\ln(\omega_m/U)$ versus $\tilde{U}$. The solid lines are the predictions of (46), with the constant $c(N)$ fit to the numerically determined points (and found in practice to depend rather slowly on $N$). The agreement between the numerics and (46) is excellent.

$\omega_m \sim c(N)U \exp\left(-\frac{\pi U}{8N\Delta_0}\right)$, (46)

The small spectral ‘glitch’ (seen e.g. at $\tilde{\omega} \approx 10^{-1}$ in the solid line of figure 3) is a well-known artefact of the RPA ladder sum used within the LMA. While its effect on the spectrum is slight, it can be removed if so desired by the procedure described in [45].
for logarithmic scale, together with comparison (dotted lines) to the scaling form over the $ω/ω_N$ of $ω$ versus $m$.

Having identified the Kondo scale $ω_m$, we return to the issue of spectral scaling in more detail. Figure 5 shows explicitly the spectral collapse of the $SU(4)$ Kondo resonance as a function of $ω/ω_m$, for $U = 30, 40$ and 50 (all collapse perfectly to the scaling form over the $σ/ω_m$ range shown). For comparison, the LMA $SU(2)$ scaling spectrum [19, 45] is also shown, dotted line. The figure highlights the clear $N$-dependence of the $SU(2N)$ scaling spectrum, alluded to in section 4.1. The $SU(4)$ Kondo scaling resonance is seen to be somewhat narrower in form compared to its $SU(2)$ counterpart (although it is of course broader on an ‘absolute’ $σ$-scale, reflecting as discussed above that $ω_m(N = 4) ≫ ω_m(N = 2)$ for given $U$).

The scaling behaviour of the tails themselves is considered in figure 5(b). Here we show the universal forms of $πΔ_0D(ω)$ versus $ω/ω_m$ for the $SU(2)$, $SU(4)$ and $SU(6)$ cases, plotted with logarithmic axes to highlight the $N$-dependence of the spectral tails (the corresponding linear plot is shown in figure 5(c) for comparison). As $N$ increases, the tails are seen to decay more rapidly, although the form of the decay is evidently similar in each case.

This behaviour of the scaling spectrum, for $|ω|/ω_m ≫ 1$ and arbitrary $N$, can in fact be obtained in closed form, following closely the approach of [45]. Details of the calculation are given in appendix C: the final result is found to be

$$\pi Δ_0D(ω) \sim \frac{1}{2} \left( \frac{4N}{\ln (|ω|)^2 + 1} \right)$$

where $ω = ω/ω_m$. The asymptotic results for $N = 1, 2$ and 3 are shown in figure 5 as dotted lines, and are seen to agree excellently with the numerical curves for $|ω| ≫ 5$.

4.4. Comparison with NRG

We conclude our discussion of the $SU(2N)$ AIM by comparing the scaling spectra obtained above with essentially exact numerics from NRG [7, 8, 39, 63, 64], employing the full density matrix formulation of the method [63, 64], and with the self-energy determined directly [65]. At the time of writing, available computational power renders it impossible to obtain accurate NRG results for models with $N > 2$. Nevertheless, the LMA scaling spectrum for $N = 1$ is known already to agree very well with the NRG result [45], and here we show that the same is true for $N = 2$. As such, it is difficult to imagine the situation changing significantly for higher values of $N$.

Figure 6 compares the LMA scaling spectra for the $SU(2)$ and $SU(4)$ models (solid and dashed lines respectively) with those of NRG (dotted); shown as a function of $ω/ω_K$ with $ω_K$ defined by $πΔ_0D(ω_K) = 1/2$. Note that both $SU(4)$ spectra have been shifted upwards by 0.4 for clarity.

The agreement between the LMA and NRG scaling spectra is striking. The tails of the spectra for $|ω|/ω_K ≫ 1$ are essentially indistinguishable, suggesting that the asymptotic form in (47) may be exact. While the LMA and NRG spectra begin to deviate when $ω/ω_K$ is reduced (in part due to the RPA ‘glitch’ in the LMA spectra), symmetry restoration ensures the correct spectral pinning required by the Friedel sum rule, namely $πΔ_0D(ω = 0) = 1$, and hence the LMA and NRG spectra once again coincide as $ω → 0$.

5. Away from $SU(2N)$ symmetry

The discussion of sections 2–4 has naturally focused on the fully symmetric $SU(2N)$ Hamiltonian of (4). We now comment briefly on the effect of an anisotropic interaction between particles on the impurity. Taking (4) and replacing the coefficient of the final term by $U'/2$, namely

$$\hat{H} = \sum_j \left[ \sum_{k,σ} ε_k c_{kjσ}^\dagger c_{kiσ} + ε \hat{n}_i + U\hat{n}_i\hat{n}_i \right]
+ V \sum_{k,σ}(c_{kiσ}^\dagger f_{iσ} + \text{h.c.}) + \frac{U'}{2} \sum_{i,j} \hat{n}_i\hat{n}_j,$$
leads to a model of capacitively coupled $SU(2)$ AIMS with distinct onsite and intersite interactions, $U$ and $U'$ respectively [40–42]. For $U' \neq U$, the Hamiltonian has $SU(2)^N$ symmetry rather than the full $SU(2N)$ symmetry of the parent model. We shall focus here on the case $U' < U$ for simplicity.

The consequence of explicit symmetry-breaking in the Hamiltonian (48) is that for $U' < U$, the lowest-energy broken-symmetry mean-field solutions are now only those with uniform charge on all sites $i$ and broken spin-symmetry. On repeating the arguments of section 3.3 however, the structure of the MF Green function for $U' < U$ remains precisely that given in (26) and (27). The LMA can then be set up in the same way as described in section 3.4, taking care now to distinguish between $U$ and $U'$ vertices in figure 1. Since each term in figure 1(b) involves repeated interactions between $(i, \sigma)$ and the same $(j, \sigma')$, each term involves either $U$ or $U'$ vertices, not a mixture of the two. It is then readily shown that the LMA self-energy of (38) generalizes to

$$\Sigma_{i\sigma}(\omega) = \Sigma_{i\sigma}(\omega; U) + (N-1)[\Sigma_{i\sigma}(\omega; U') + \Sigma_{i\sigma}(\omega; U')]$$

(49)

where the $U$ or $U'$ vertices appearing in the various terms are labelled explicitly. For the reasons discussed in appendix B, the contribution of the $\Sigma_{i\sigma}(\omega; U')$ term in (49) is negligible when $U' \gg 1$. The two remaining terms by contrast both involve the transverse spin polarization propagator $\Pi(\omega)$, the resonance in which generates the low-energy Kondo scale within the LMA (see section 4.2). Neither term can therefore be neglected a priori. And since each is calculated with different interaction vertices, there are now two distinct energy scales in the problem when $U' \neq U$.

We have studied the ensuing $U' < U$ behaviour in detail, both numerically and where possible analytically. The typical behaviour of the two energy scales is shown for the illustrative case of $SU(4)$ ($N = 2$) with $\tilde{U} = 20$ in figure 7(a). We denote by $\omega_m$ the scale that enters the first term in (49) (arising from spin-flip excitations on the same site), shown in figure 7(a), while the other scale (due to spin-flip excitations involving distinct sites) is labelled $\omega_m'$ and also shown in the figure.

The two-scale behaviour is in good agreement with NRG results [40, 41]. The low-energy scale $\omega_m$ deviates very little from its $U = 0$ $SU(2)$ behaviour ($\propto \exp(-\pi U/8)$, (46) with $N = 1$) on initially increasing $U'/U$. Only when $U - U'$ itself becomes of the order of the $SU(4)$ Kondo scale does the scale rapidly cross over to the $U' = U$ behaviour $[\omega_m \propto \exp(-\pi U/16)]$ discussed in section 4.2 [40]. The higher-energy scale $\omega_m'$ is typically of order $U - U'$, and arises in the NRG as a crossover scale from the $SU(4)$ local moment (LM) fixed point to the $SU(2)$ LM fixed point [41]. And as $U' \rightarrow U$, figure 7(a) shows that $\omega_m' \rightarrow \omega_m$, reflecting the natural existence of only a single scale at the $SU(4)$ point $U' = U$.

![Figure 6](image1.png)

**Figure 6.** Comparison between LMA and NRG scaling spectra. Solid and dashed lines show respectively the LMA $SU(2)$ and $SU(4)$ scaling spectra, while the dotted lines are corresponding NRG results. The Kondo scale $\omega_m$ is defined by $\pi \Delta_0 D(\omega_m) = 1/2$. Note that the $SU(4)$ spectra have been shifted upward by 0.4 for clarity.

![Figure 7](image2.png)

**Figure 7.** Results away from $SU(2N)$ symmetry. (a) The behaviour of the two energy scales $\omega_m/\Delta_0$ (solid line) and $\omega_m'/\Delta_0$ (dashed line), as a function of $U'/U$, for fixed $\tilde{U} = 20$ and $N = 2$. (b) The universal single-particle spectrum $\pi \Delta_0 D(\omega)$ calculated close to the $SU(4)$-symmetric point, with $\alpha = (U - U')/\omega_m(U' = U) = 5$ (solid line). Also shown is the pure $SU(2)$ scaling spectrum (dashed line), and the pure $SU(4)$ scaling spectrum (dotted).
From our current perspective the most interesting physics is seen when the two scales are comparable; i.e., when $U - U' \sim \mathcal{O}(\alpha_m)$. To express this more precisely, we define the quantity

$$\alpha = \frac{U - U'}{\alpha_m(U' = U)}$$

(50)

with $\alpha_m(U' = U)$ the $SU(4)$ Kondo scale arising when $U' = U$. For fixed $N$ we find that when $\bar{U} \gg 1$, the ratios $\alpha_m/\alpha_m$ and $\alpha_m/\alpha_m(U' = U) = \gamma(\alpha)$ are universal functions of $\alpha$ for any finite $\alpha$. As a result, spectra for fixed $\alpha > 0$ show universal scaling collapse in the same way as seen previously for $\alpha = 0$ (section 4.3).

We show a representative example of one of these scaling spectra in figure 7(b), for the case $\alpha = 5$ (solid line). Note that in addition to the spectral 'glitch' at $\omega = \omega_m$, there is now another at $\omega = \omega_m' \simeq 25\omega_m$. Both are a simple consequence of the RPA self-energy approximation (and can be removed if so desired [45]), although in this case they serve a useful purpose in marking out the low-energy scales of the model. Figure 7(b) shows in addition the universal $SU(2)$ scaling spectrum as a function of $\omega/\omega_m$ (dashed line), and the universal $SU(4)$ spectrum also as a function of $\omega/\omega_m = \omega/\gamma(\alpha)\omega_m(U' = U)$ (dotted line). Just as found in NRG studies [42], we see immediately that the $U' < U$ scaling spectrum shows a clear crossover from universal $SU(4)$ behaviour when $|\omega| \gg \omega_m'$, to universal $SU(2)$ behaviour when $|\omega| \ll \omega_m'$, reflecting the different effective low-energy models that arise on the two distinct energy scales [40, 42].

6. Conclusions

We have considered here a local moment approach to single-particle dynamics of the orbitally degenerate $SU(2N)$ Anderson model; focusing explicitly on the particle–hole asymmetric case for arbitrary $N$, where the impurity is occupied by $N$ electrons—a tangible example being the middle of the 2-electron Coulomb blockade valley in a carbon nanotube quantum dot with $SU(4)$ symmetry [21–24]. The resultant LMA is a natural generalization of the technique developed previously for $SU(2)$ quantum impurity models; and is seen to provide a rather good description of the dynamics of the $SU(2N)$ Anderson model, recovering the correct exponential vanishing of the Kondo scale for $\bar{U} \gg 1$ and its associated universality, the broadening of the Hubbard satellites by many-body scattering processes, and a very good description of the single-particle scaling spectrum as judged by direct comparison to NRG results for the $SU(4)$ and $SU(2)$ models.

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Appendix A. Closed form of $0\Pi^{\sigma\sigma}(\omega)$

The longitudinal spin polarization bubble $0\Pi^{\sigma\sigma}(\omega)$ is calculated from (30) and (41). We find

$$\pi \Delta_0 \Re 0\Pi^{\sigma\sigma}(\tilde{\omega}) = \frac{4\sigma f(\tilde{\omega}) + \omega g(\tilde{\omega})}{2\hbar(\tilde{\omega})}$$

(A.1)

$$\sgn(\omega)\pi \Delta_0 \Im 0\Pi^{\sigma\sigma}(\tilde{\omega}) = \frac{\sigma |\tilde{\omega}| + g(\tilde{\omega})}{\hbar(\tilde{\omega})}$$

(A.2)

where

$$f(\tilde{\omega}) = \tan^{-1}(\tilde{\omega} + \tilde{x}) - \tan^{-1}(\tilde{\omega} - \tilde{x})$$

(A.3)

$$g(\tilde{\omega}) = \ln\left\{\frac{\sigma + \tilde{x}^2}{\tilde{\omega}^2 + \tilde{x}^2 + 1}\right\}$$

(A.4)

$$h(\tilde{\omega}) = \tilde{\omega}(\tilde{\omega}^2 + 1).$$

(A.5)

Appendix B. Derivation of (46)

The derivation of (46) for arbitrary $N$ is closely analogous to that of the $N = 1$ case [19]. We begin by noting from (20) that the MF moment $|\bar{m}_0| \to 1$ when $\bar{U} \to \infty$. Since stability of the RPA polarization propagators requires (43), symmetry restoration takes place at an $x \simeq U/2$ and thus, from (37), $|\bar{\mu}| \to 1$ also. Hence the symmetry restoration condition, (45), becomes

$$\Sigma_{1+}(\omega = 0) \lesssim \frac{U}{2},$$

(B.1)

or, equivalently,

$$\frac{N\Sigma_{1-}^{-}(\omega = 0) + (N - 1) \Sigma_{1+}^{++}(\omega = 0) \lesssim \frac{U}{2}}{U}$$

(B.2)

from (38).

The first term on the left-hand side of (B.2) is simply $N$ times the LMA self-energy for the $SU(2)$ AIM, the asymptotic $\bar{U} \gg 1$ behaviour of which has already been examined in detail [19]. The key is to note that the RPA $\Pi^{++}(\omega)$ is dominated by a sharp resonance at $\omega = \omega_m$, which as $\bar{U} \to \infty$ tends asymptotically to a delta-function, leading to [19]

$$\Sigma_{1+}^{++}(\omega = 0) \lesssim \frac{4\Delta_0}{\pi} \ln\left(\frac{U}{\alpha_m}\right).$$

(B.3)

The second term on the left-hand side of (B.2) can be analysed in a similar fashion, starting from the closed form of $0\Pi^{++}(\omega)$ in appendix A, and using (40) to obtain the RPA longitudinal spin polarization propagator $\Pi^{++}(\omega)$. Crucially, in marked contrast to the $\Pi^{--}(\omega)$ discussed above, we find that no sharp resonance develops in the RPA $\Pi^{++}(\omega)$ when $\bar{U} \gg 1$. The numerically determined $\Sigma_{1-}^{-}(\omega = 0)$ thus turns out to be $\mathcal{O}(U^{-1})$, and hence in the limit $\bar{U} \gg 1$, $\Sigma_{1+}^{++}(\omega = 0)$ can be neglected in comparison to the logarithmically diverging $\Sigma_{1+}^{++}(\omega = 0)$ of (B.3). Combining (B.2) and (B.3) then leads to (46) straightforwardly.
Appendix C. Derivation of (47)

The tails of the SU(2N) scaling spectra can be obtained in closed form within the LMA. We define $\tilde{\omega} = \omega/\omega_m$ and consider the behaviour of the LMA self-energies for fixed $\tilde{\omega}$ in the formal scaling limit $\omega_m \rightarrow 0$. Let us rewrite (38) explicitly as

$$\Sigma_{i\sigma} (\tilde{\omega} \omega_m) = N \Sigma_{i\sigma}^{0} (\tilde{\omega} \omega_m) + (N - 1) \Sigma_{i\sigma}^{0} (\tilde{\omega} \omega_m). \quad (C.1)$$

As described in appendix B the imaginary part of the longitudinal spin polarization propagator, $\text{Im} \Pi^{\sigma\sigma} (\omega)$, contains only broad resonances at $\omega \approx \mathcal{O}(U)$ rather than the emerging delta-function at $\omega = \omega_m$ in $\text{Im} \Pi^{\sigma\sigma} (\omega)$. This means that $\Sigma_{i\sigma}^{0} (\omega)$ is non-universal at low frequencies and hence, in the limit $\omega_m \rightarrow 0$ for fixed $\tilde{\omega}$, we can write $\Sigma_{i\sigma}^{0} (\tilde{\omega} \omega_m) = \Sigma_{i\sigma}^{0} (0)$. But since $\Sigma_{i\sigma}^{0} (0) = \mathcal{O}(U^{-1})$ (as in appendix B), it vanishes in the formal scaling limit $\tilde{U} \rightarrow \infty$. Hence (C.1) reduces simply to

$$\Sigma_{i\sigma} (\tilde{\omega} \omega_m) \approx N \Sigma_{i\sigma}^{0} (\tilde{\omega} \omega_m). \quad (C.2)$$

Moreover, $\Sigma_{i\sigma}^{0} (\tilde{\omega} \omega_m)$ is precisely the function already analysed to obtain the scaling spectrum of the SU(2) AIM [45]. All the analysis of [45] can thus be generalized to the SU(2N) model quite straightforwardly. The upshot is that equation (5.1) of [45] must simply be multiplied by $N$ (reflecting the additional factor of $N$ in (C.2)) which, by analogy to equation (5.2) of [45], gives the full LMA scaling spectrum for the SU(2N) AIM as

$$\pi \Delta_{0} E(D) = \frac{1}{2} \left[ \frac{1}{[\tilde{\omega} + 1]^{2} + 1} + \frac{1}{[\tilde{\omega} - 1]^{2} + 1} + 4N\theta (\tilde{\omega} - 1) \right]$$

for $\tilde{\omega} > 0$ (the corresponding negative-frequency spectrum follows from particle–hole symmetry, i.e. $D(-\omega) = D(\omega)$). In the limit $|\tilde{\omega}| \gg 1$, (47) results.

References