In what state of equilibrium can a quantum system be? Does this state have universal properties and what are the conditions for its emergence? These questions are not new, dating even from the very birth of quantum theory \cite{1} and are surprisingly open\cite{2,3}. Indeed, the foundations of statistical physics still rely today on a static Bayesian point of view assuming the equiprobability of the accessible states defining the microcanonical ensemble. Assuming temperature and chemical potential can be defined then the canonical and grand canonical ensembles can be derived, allowing to calculate all relevant macroscopic quantities in the thermodynamical limit \cite{4–6}. In order to link theoretical predictions calculated with averages over these ensembles to experimental quantities measured on a single system, an assumption of ergodicity is made. Despite being broadly accepted, this assumption is not justified in a satisfactory manner (see, e.g., the discussion in Ref. \cite{7}). Triggered by recent progress in the quantum engineering of mesoscopic systems \cite{8,9}, some theoretical progress has been achieved for attempting to explain thermodynamical equilibrium with a purely quantum point of view.

From the early work of von Neumann on quantum ergodicity \cite{10,11}, most theoretical studies aiming at understanding thermalisation as a quantum and universal \cite{11} process have focused on looking for signatures of thermalisation on physical observables of large quantum systems \cite{12,15}, for instance with the Eigenstate Thermalisation Hypothesis (ETH) surmise \cite{10,13}. Instead of observables, one can also focus on the state of a system embedded in a larger one for which a “canonical typicality” property has been established: the overwhelming majority of pure quantum states of the composite system are locally\cite{14} canonical \cite{20,22}. This static “typicality” has been extended to the dynamics of embedded quantum systems (two-level \cite{23}, four-level \cite{24} and arbitrary \cite{25} quantum systems). We apply here this “dynamical typicality” property in order to calculate analytically and with full generality the stationary state of an embedded quantum system at long time. We find a new thermodynamical ensemble of purely quantum origin characterizing this state. This ensemble captures the microcanonical and the canonical ensembles as particular cases, and as such provides a quantum explanation for the Gibbs distribution.

We consider an arbitrary quantum system coupled to a large arbitrary quantum environment through a random interaction. We emphasize the fact that the initial state of this composite system can be chosen arbitrarily, in particular the environment does not have to be in thermal equilibrium initially nor the full composite system in the microcanonical situation. Dynamical typicality \cite{24} states that for almost all interaction Hamiltonian\cite{26} the reduced density matrix of the system has a self-averaging property in the large environment limit \cite{27}, in other words, it follows a universal dynamics. Despite this does not imply a priori equilibration, since it can be consistent with sustained oscillations and revivals\cite{28}, this property has a very practical consequence. It allows to perform non perturbative analytical calculations with full generality, i.e. for arbitrary system, environment, and global initial state, by justifying rigorously an averaging procedure over some randomness introduced only at the level of the interaction Hamiltonian. We apply this calculation framework here to study the state of the system at long but finite times, i.e. smaller than any recurrence time. Postponing all questions regarding the out of equilibrium dynamics to a further publication \cite{29}, we show that: if the system converges towards a stationary state, then this state is characterized by a new quantum partition function which can be calculated. This partition function relies on an average transition probability between states involving some purely quantum quantities: the fourth order moments of the overlap coefficients between eigenvectors of a bare and a dressed Hamiltonian. We calculate this transition probability for several classes of random interactions. Then we calculate the probabilities of occupation of the states of the system a find a new thermodynamical ensemble more general than the microcanonical one.
I. MODEL SETUP

The setup is identical to [23]: we consider a system $S$ in contact with an environment $E$, writing $\mathcal{H}_s, \mathcal{H}_e$ for their respective Hilbert spaces. The total system $S + E$ is closed and its Hilbert space is the tensor product $\mathcal{H} = \mathcal{H}_s \otimes \mathcal{H}_e$ (with dimension $N = \dim \mathcal{H}_e \dim \mathcal{H}_s$). The total or dressed Hamiltonian $\hat{H}$ is the sum $\hat{H} = \hat{H}_s + \hat{H}_e + \hat{W}$ where $\hat{W}$ is an interaction term. Eigenvectors of the “bare” Hamiltonian $\hat{H}_s + \hat{H}_e$ are written as $|\phi_n\rangle$ and are tensor products of eigenvectors $|\epsilon_n\rangle$ of $\hat{H}_e$ and eigenvectors $|\epsilon_c\rangle$ of $\hat{H}_c$, with the eigenenergies $\epsilon_n = \epsilon_s + \epsilon_c$. We write $|\psi_i\rangle$ for the dressed eigenvectors and $\{\lambda_i\}_i$; the set of associated dressed eigenvalues. The state of $S + E$ is described by a density matrix $\rho(t)$ which follows the well known relation

$$\rho(t) = \hat{U}_t \rho(0) \hat{U}_t^\dagger \quad \text{with} \quad \hat{U}_t = e^{-\frac{i}{\hbar} \hat{H} t}.$$ 

The state of the subsystem $S$ is described by a reduced density matrix: $\rho_s(t) = \text{Tr}_E \rho(t)$, $\text{Tr}_E$ being the partial trace with respect to the environment. Decomposing the initial state $\rho(0)$ on the bare eigenbasis $\{|\phi_1\rangle,...,|\phi_N\rangle\}$ and using linearity, we consider the matrix elements expanding the evolution operator $\hat{\rho}$ density matrix:

$$S$$

To calculate the expression in Eq. (1), one needs an analytical formula for the overlap coefficients $\langle \psi_i | \phi_n \rangle$ and the dressed eigenvalues $\lambda_i$, which are quantities usually accessible in a perturbative framework only. In this Letter, we use a statistical method for calculating these quantities in a non perturbative setting and for arbitrary system and environment.

The method relies on the hypothesis assumed for the interaction Hamiltonian: we introduce deliberately some randomness in and only in the interaction $\hat{W}$ in order to perform calculations, knowing that this randomness actually will not matter in the large dimensionality limit (dim $\mathcal{H}_e \to \infty$) due to the typicality of the dynamics [23]. This randomness should be compatible with some macroscopic constraints: $\hat{W}$ “centered” i.e. $\text{Tr}(\hat{W}) = 0$ and with fixed spectrum variance $\sigma^2_\rho = \text{Tr}(\hat{\rho}(\hat{W}^\dagger \hat{W})/N$ independent of $N$. Then regarding the symmetry class of the randomness, we will assume $\hat{W}$ to be either a Wigner band random matrix (WBRM) [20] or a randomly rotated matrix (RRM i.e. of the type $\hat{U} \hat{Q} \hat{U}^\dagger$ with $\hat{Q}$ real diagonal fixed and $\hat{U}$ unitary or orthogonal Haar distributed). The WBRM ensembles are convenient for modeling interactions in heavy atoms and nuclei [31-33]. The sparsity of WBRM comes from the finite energy range of the interaction. On the other hand, RRM ensembles are dense, which contradicts the a priori two body nature of the interaction, but provides a convenient way for modeling the local spectral statistics of more physical interaction Hamiltonians [33-37].

II. TYPICAL DYNAMICS

We now focus on the reduced density matrix: $\rho_s(t) = \text{Tr}_E \rho(t)$ and consider it as a function of the interaction $\hat{W}$, keeping all other parameters constant (time, spectra of $S$ and $E$, initial state). This function exhibits a generalized central limit theorem phenomenon known as the concentration of measure [23].

A. Concentration of measure

This phenomenon can be described informally as follows: a numerical function which depends on many independent random variables in a balanced way (i.e. there is no outliers on which this function depends) is very close to its mean value almost everywhere. On the quantitative side, this phenomenon can be characterized rigorously with the following upper bound on the variance of this function away from its mean behavior [25]:

$$\sigma^2_\rho = \mathbb{E}[(\rho_s(t) - \mathbb{E}[\rho(t)])^2] \leq \frac{4 \sigma^2_\rho}{\hbar^2} \frac{1}{\dim \mathcal{H}_e},$$

where $\mathbb{E}$ is the average over the set of interaction Hamiltonians considered (WBRM and RRM) and $||\hat{A}||^2 = \text{Tr}(\hat{AA}^\dagger)$. As $\dim \mathcal{H}_e \to \infty$, $\sigma^2_\rho \to 0$ and consequently $\rho_s(t)$ is getting very close to its mean value which provides the typical dynamics. We can thus compute an approximate $\rho_s(t)$ simply by averaging: $\rho_s(t) = \text{Tr}_E(\rho(t)) \approx \mathbb{E}[\text{Tr}_E(\rho(t))] = \text{Tr}_E(\mathbb{E}[\rho(t)])$. We are led to consider the average of Eq. (1).

We will now focus specifically on the stationary regime at long times. Under the hypothesis assumed on the statistics of the interaction (WBRM and RRM ensembles) the dressed eigenvalues $\{\lambda_1,...,\lambda_N\}$ undergo level repulsion and, as such, are non degenerate. This implies that the time independent terms are provided by the case $i = j$ in the summation in Eq. (1) averaged over $\hat{W}$:

$$\sum_i \mathbb{E}[\langle \phi_n | \psi_i \rangle \langle \phi_i | \phi_m \rangle \langle \phi_p | \psi_i \rangle \langle \psi_i | \phi_q \rangle].$$

The time dependent regime (given by the summation over $i$ and $j$ such that $i \neq j$) is outside the scope of this article. We will assume this regime to be damped (see [34] for $\hat{W}$ in the WBRM ensemble), without revivals [28] at least on the largest time scale of this model ($1/D$ where $D$ is the mean level spacing of the dressed Hamiltonian) such that considering a stationary regime is meaningful over this time scale.
We first single out the non zero cases for the fourth order moments of the overlap coefficients:  
\[ E[|\langle \phi_n | \psi_i \rangle|^2 |\langle \phi_m | \psi_i \rangle|^2] \]  
which are when \( n = m \) and \( p = 0 \) or when \( n = q \) and \( m = p \) [40]. The former case is involved in the asymptotic value of the off-diagonal terms of \( g_s(t) \) i.e. the quantum coherences of the state of \( S \), which can be shown to be zero as expected in the limit \( t \to \infty \) [41]. In the following, we focus on the later case \( (n = q \text{ and } m = p) \) which governs the dynamics of the diagonal terms of \( g(t) \) and \( g_s(t) \), i.e. the probabilities of occupation.

### B. Average transition probability

We define from Eq. 2 with \( n = q \) and \( m = p \), an average transition probability \( \bar{p}_{m \to n} \) from an initial state \( |\phi_m \rangle \) at \( t = 0 \) to a final state \( |\phi_n \rangle \) at \( t \to \infty \):

\[
\bar{p}_{m \to n} = \sum_i E[|\langle \phi_n | \psi_i \rangle|^2 |\langle \phi_m | \psi_i \rangle|^2].
\]  
(3)

Such sum provides quantitatively how \( |\phi_n \rangle \) is accessible from \( |\phi_m \rangle \) and has been considered, e.g., numerically in the context of random two body interactions (TBRI) ensembles [41] and analytically for some specific systems: quantum walkers [42–44]. The particular case \( m = n \) provides the return probability whose reciprocal 1/\( \bar{p}_{m \to n} \) is the so-called purity [45–48]. The leading order of \( \bar{p}_{m \to n} \) is given by \( \sum_i E[|\langle \phi_n | \psi_i \rangle|^2] E[|\langle \phi_m | \psi_i \rangle|^2] \) and involves the second order moment of the overlaps \( E[|\langle \phi_n | \psi_i \rangle|^2] \). This quantity, multiplied by \( \rho \), is called the local density of states (LDOS) and quantifies how much a bare eigenvector is delocalized or hybridized with the dressed eigenbasis and has already been considered in various contexts (nuclear physics [49], molecular physics [50, 51], atomic physics [52], thermalisation [12], quantum chaos [31], financial data analysis [53, 54], see also the review in [33]) for various cases of \( H_0 \) and \( W \). It has the following typical shape:

\[
E[|\langle \phi_n | \psi_i \rangle|^2] \approx \frac{f(\lambda_i - \epsilon_n)}{\int \rho_{e+}(\epsilon) f(\lambda_i - \epsilon) d\epsilon}
\]  
(4)

where \( \rho_{e+} \) is the bare density of states, \( \lambda_i \) is the mean of the dressed eigenvalue \( \lambda \) and \( f \) is a function peaked around zero with a typical width \( \Gamma \). The denominator is here for the purpose of normalisation. For most models of \( H_0 \) and \( W \), the function \( f \) is a Lorentzian reminiscent of the Breit-Wigner law with a generalized Fermi Golden rule rate \( f = \frac{\pi}{\sqrt{\gamma^2 - \rho}^2} \). This provides the main result of this paper: for an initial state \( \phi(0) = |\phi_m\rangle \langle \phi_m| \), the long time stationary state of \( S \) is distributed according to

\[
p_{\epsilon_s} = \lim_{t \to \infty} \langle \epsilon_s | g_s(t) | \epsilon_s \rangle \approx \int \frac{\rho_{e+}(\epsilon) g(\epsilon_m - \epsilon_s) d\epsilon}{\int \rho_{e+}(\epsilon) g(\epsilon_m - \epsilon) d\epsilon}.
\]  
(6)

The denominator is the convolution of the bare density of states by the transition probability \( g \) which provides the effective number of bare states accessible from the initial \( |\phi_m\rangle \). Such a quantity \( \rho_{e+}(\epsilon) g(\epsilon_m - \epsilon) d\epsilon \) enforces the normalization condition and can be considered as a new partition function. The numerator is the convolution of the environment density of states by the transition probability \( g \) and provides the effective number of accessible states such that \( S \) is in the state of energy \( \epsilon_s \). The probability of occupancy is the ratio of these two numbers. Let us now consider the case of intermediate coupling.

### III. MAIN RESULT

#### A. General case

In order to perform the partial trace and get \( g_s(t) \), we recall the final state \( |\phi_n\rangle = |\epsilon_s\rangle |\epsilon_e\rangle \) and sum Eq. 5 over \( \epsilon_s \) using a continuous approximation: \( \text{Tr}_{e} = \sum_{\epsilon_s} \leftrightarrow \int d\epsilon \rho_{e}(\epsilon) \). This provides the main result of this paper: for an initial state \( \phi(0) = |\phi_m\rangle \langle \phi_m| \), the long time stationary state of \( S \) is distributed according to
B. Intermediate coupling case

A temperature can be defined by $\beta = \frac{1}{kT} = \frac{d \ln \rho_{e}}{d \epsilon}$. Assuming a good decoupling between the micro ($D = 1/\rho$), meso ($\Gamma'$) and macro ($kT$) energy scales: $D \ll \Gamma' \ll kT$, and considering all energies $\epsilon_{s}, \epsilon_{m}$ to be inside the bulk of the spectrum, then the function $g$ in Eq. (4) can be approximated by a Dirac function which is “sampling” $\rho_{e}(\epsilon)$ at $\epsilon_{e} = \epsilon_{m} - \epsilon_{s}$ and simplifying Eq. (6) for

$$p_{\epsilon_{s}} \approx \frac{\rho_{e}(\epsilon_{m} - \epsilon_{s})}{\rho_{e}(\epsilon_{m})} \approx \frac{\rho_{e}(\epsilon_{m} - \epsilon_{s})}{\rho_{e}(\epsilon_{m})} \approx e^{-\beta \epsilon_{s}} Z_{\beta}$$

with $Z_{\beta} = \sum_{\epsilon_{s}} e^{-\beta \epsilon_{s}}$ the canonical partition function. In other words, the Boltzmann distribution is a particular case of the more general distribution provided by Eq. (6) whose origin is quantum.

C. Strong coupling case

If the coupling is strong enough that $\Gamma' \gtrsim kT$ then the transition probability $g$ cannot be approximated by a Dirac function and its finite width must be taken into account in the convolution in Eq. (6). From this convolution effect, one should expect a decrease of contrast in the probability distribution of $S$ when the interaction strength is increased: the equilibrium probability then undergoes a continuous crossover from the local microcanonical ensemble prediction we described earlier (i.e. equiprobability over a small energy shell of accessible states around initial energy) to a global microcanonical ensemble prediction (i.e. all bare state are accessible and equiprobable). The convolution in Eq. (6) can be done analytically e.g. when $\rho_{e}$ is Gaussian and $g$ is Lorentzian: one obtains the Voigt distribution, relevant in atomic spectroscopy when a natural linewidth is broadened by the Doppler effect. We check numerically these predictions on Fig. 4 and find a satisfactory agreement.

Finally, we stress that the above results are valid for an initial state $|\phi_{m}\rangle \otimes |\phi_{p}\rangle = |\epsilon_{s}\rangle \otimes |\epsilon_{c}\rangle |\epsilon_{c}\rangle$ and can be extended by linearity to any initial state, pure or not: the extra diagonal terms (i.e. of the type $|\phi_{m}\rangle \langle \phi_{p}|$ with $m \neq p$) do not contribute, only the diagonal ones contribute (see [40]). Therefore the stationary state of $S$ is the weighted average of Eq. (3) by the initial energy distribution of the composite system.

IV. CONCLUSION AND SUMMARY

We showed that the stationary properties of an embedded quantum system are encoded in the geometric relation between the eigenvectors of a bare and a dressed Hamiltonian, more precisely in the fourth order moments of the overlaps between their eigenvectors. This fact
provides a purely quantum way to define a new partition function which can be calculated thanks to dynamical typicality [25]. In the intermediate coupling case $D < \Gamma' \ll kT$, this partition function simplifies to the prediction of a local microcanonical ensemble defined on a small energy window around the initial energy. In the strong coupling regime (i.e. $D \ll kT \lesssim \Gamma'$), one gets a more general ensemble which depends on the interaction strength and leads to a loss of contrast of the probabilities of occupation (i.e. a convergence towards global equiprobability). We considered here two random matrix ensembles for the interaction which have broad applicability. Our framework could be used with other interaction Hamiltonian ensembles (e.g. conserving some set of observables or enforcing the two body nature of the interaction) as soon as dynamical typicality is shown to be verified and a local density of states is available.

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[11] By universal, we mean independent of the specific Hamiltonian considered.
[19] By locally we mean after taking a partial trace to get the state of the subsystem.
[26] This “almost all” is relative to the probability measure defined on the random matrix ensemble considered.
[27] This self-averaging property was also used to investigate the local state of the microcanonical ensemble in the case of small subsystem and strong interaction [66].
[28] Because of the finite dimension of the Hilbert space, this regime might present quantum recurrences and revivals.
[29] See [23, 67] for the case of a two level system coupled to an environment through a separable random interaction.
[30] WBRM are of the type $W_{ij} = a(i-j)/bY_{ij}$ where $Y$ is a Wigner Random Matrix and $a(x)$ is a deterministic band profile, “b” being the bandwidth.