A unified approach to the derivation of work theorems for equilibrium and steady-state, classical and quantum Hamiltonian systems

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Abstract

We present a unified and simple method for deriving work theorems for classical and quantum Hamiltonian systems, both under equilibrium conditions and in a steady state. Throughout the paper, we adopt the partitioning of the total Hamiltonian into the system part, the bath part, and their coupling. We rederive many equalities which are available in the literature and obtain a number of new equalities for nonequilibrium classical and quantum systems. Our results can be useful for determining partition functions and (generalized) free energies through simulations or measurements performed on nonequilibrium systems.
I. INTRODUCTION

The fluctuation theorems allow us to rigorously relate equilibrium ensemble properties of a dynamical system with its evolution under nonequilibrium conditions, beyond the domain of validity of the linear response theory. Many recent results, concepts, and ideas are stemmed from the early landmark work by Bochkov and Kuzovlev. Among the fluctuation theorems, the Jarzynski equality (or, equivalently, the nonequilibrium work theorem) occupies a remarkable place. This equality connects the nonequilibrium work performed on a system with the ratio of the equilibrium system’s free energies. Over the past ten years, the Jarzynski equality has been extended to non-Markovian stochastic processes to quantum systems and to systems coupled to different (non-Hamiltonian) thermostats. Several important results for dissipative systems in the steady state have also been obtained.

The fluctuation theorems and Jarzynski equality have been proven for a Hamiltonian system coupled to Hamiltonian heat bath(s), see Refs. and Ref., respectively. The present paper is aimed at presenting a unified and simple method for generating various work theorems for such systems. We consider both classical and quantum systems, which can initially be prepared either under equilibrium conditions or in a steady state. Within our approach, we rederive many equalities which are available in the literature and obtain a number of new results for nonequilibrium classical and quantum systems. Our expressions can be considered as mathematical identities, since the fulfillment of the Liouville theorem is required only. They can be useful for determining various (equilibrium or steady state) partition functions and (generalized) free energies through simulations and/or measurements performed on nonequilibrium systems. The nonequilibrium partition functions can be used in much the same manner as their equilibrium counterparts. Indeed, our steady-state distributions have the generic form \( \rho_{ne} = 1/Z_{ne} \exp(\sum_n g_n G_n) \), where \( Z_{ne} \) is the non-equilibrium partition function, \( g_n \) are certain c-numbers, and \( G_n \) are the corresponding operators. If we differentiate the logarithm of the partition function with respect to the parameter \( g_n \), we obtain the expectation value of the operator \( G_n \). Doing so, we can get expectation values of the steady-state energy, entropy, particle number, etc. Caution should be exercised in quantum case, however, because operators may not commute with each other.
II. CLASSICAL SYSTEMS

Let $H(\Gamma, t)$ be the Hamiltonian (which is allowed to be explicitly time-dependent), $\Gamma$ be the corresponding phase variables, and $A(\Gamma, t), B(\Gamma, t)$ be certain non-pathological functions of the phase variables and time. Then we can write down the identity

$$A(\Gamma_0, 0)A(\Gamma_0, 0)^{-1}B(\Gamma_t, t) = B(\Gamma_t, t),$$

$\Gamma_0$ and $\Gamma_t$ being the values of the phase variables at the time moments 0 and $t$. We can integrate Eq. (1) over $\Gamma_0$ and make use of the fact that the motion of a Hamiltonian system can be regarded as a canonical transformation, for which the Liouville theorem holds: $d\Gamma_0 = d\Gamma_t$. We thus obtain

$$\int d\Gamma_0 A(\Gamma_0, 0) \left( A(\Gamma_0, 0)^{-1}B(\Gamma_t, t) \right) = \int d\Gamma_t B(\Gamma_t, t) \equiv B_t. \quad (2)$$

If we assume that $A(\Gamma_0, 0)$ is positively defined and normalized ($\int d\Gamma_0 A(\Gamma_0, 0) = 1$), we can consider $A$ as the probability density, denote the averaging

$$\int d\Gamma_0 A(\Gamma_0, 0)... \equiv \langle ... \rangle_A \quad (3)$$

and rewrite Eq. (2) as

$$\langle A(\Gamma_0, 0)^{-1}B(\Gamma_t, t) \rangle_A = B_t. \quad (4)$$

A. Systems at equilibrium

The proof of Eq. (4) given above is very similar to that of the Jarzynski equality for Hamiltonian systems. However, Eq. (4) contains the Jarzynski equality and much more. Indeed, let both $A$ and $B$ be the instantaneous Gibbs distributions:

$$A(\Gamma_0, t) = \rho_0(\Gamma_0) = Z_0^{-1} \exp(-\beta H(\Gamma_0, 0)), \quad B(\Gamma_t, t) = \rho_t(\Gamma_t) = Z_t^{-1} \exp(-\beta H(\Gamma_t, t)). \quad (5)$$

Here $H(\Gamma_t, t)$ is a time-dependent Hamiltonian,

$$Z_0 = \int d\Gamma_0 \exp(-\beta H(\Gamma_0, 0)), \quad Z_t = \int d\Gamma_t \exp(-\beta H(\Gamma_t, t)) \quad (6)$$

are the corresponding partition functions and $\beta = 1/(k_B T)$, $k_B$ being the Boltzmann constant and $T$ being the temperature. Plugging Eqs. (5) into our starting formula (4) we get

$$\langle \exp(-\beta(H(\Gamma_t, t) - H(\Gamma_0, 0))) \rangle_{\rho_0} = Z_t/Z_0. \quad (7)$$
The time derivative of any function $C(\Gamma_t, t)$ is determined by the following expression:

$$\frac{d}{dt} C(\Gamma_t, t) = \frac{\partial}{\partial t} C(\Gamma_t, t) + \{C(\Gamma_t, t), H(\Gamma_t, t)\},$$

(8)

{$\{\}$} being the Poisson bracket. Thus

$$C(\Gamma_t, t) - C(\Gamma_\tau, \tau) \equiv \int_\tau^t d\tau' \frac{\partial'}{\partial \tau'} C(\Gamma_{\tau'}, \tau') = \int_\tau^t d\tau' \frac{\partial'}{\partial \tau'} C(\Gamma_{\tau'}, \tau') + \int_\tau^t d\tau' \frac{\partial'}{\partial \tau'} C(\Gamma_{\tau'}, \tau').$$

(9)

We partition the total Hamiltonian into the system Hamiltonian, the bath Hamiltonian, and their coupling:

$$H(\Gamma_t, t) = H_S(x_t, t) + H_B(X_t) + H_{SB}(x_t, X_t).$$

(10)

Here \(x_t\) and \(X_t\) are the phase variables specifying the system and the bath, and the system Hamiltonian only is allowed to be explicitly time-dependent. Plugging $H(\Gamma_t, t)$ (10) into identity (9) and making use of the fact that \(\{H(\Gamma_t, t), H(\Gamma_t, t)\} \equiv 0\), we can write

$$H(\Gamma_t, t) - H(\Gamma_0, 0) = \int_0^t dt' \frac{\partial}{\partial t'} H(\Gamma_{\tau'}, \tau') = \int_0^t dt' \frac{\partial}{\partial t'} H_S(x_{\tau'}, \tau') \equiv W,$$

(11)

\(W\) being the work performed on the system. Thus Eqs. (7) and (11) yield the Jarzynski formula [11, 35]

$$\langle \exp(-\beta W) \rangle_{\rho_0} = \frac{Z_t}{Z_0}.$$  

(12)

Eq. (11) can be applied to more complicated situations. Let us assume that the system and the bath are initially prepared at different temperatures \(T_S (\beta_S = 1/(k_B T_S))\) and \(T\), respectively. We can take \(A\) and \(B\) to be the corresponding nonequilibrium distributions

$$A(\Gamma_0, 0) = \rho_{\beta_0}(\Gamma_0) = Z_{\beta_0}^{-1} \exp(-\beta H(\Gamma_0, 0) - (\beta_S - \beta) H_S(x_0, 0)),$$

$$B(\Gamma_t, t) = \rho_{\beta_t}(\Gamma_t) = Z_{\beta_t}^{-1} \exp(-\beta H(\Gamma_t, t) - (\beta_S - \beta) H_S(x_t, t)),$$

(13)

\(Z_{\beta_0}\) and \(Z_{\beta_t}\) being the corresponding partition functions. Inserting these formulas into Eq. (11), we obtain

$$\langle \exp(-\beta_S W - (\beta_S - \beta) Q) \rangle_{\rho_{\beta_0}} = \frac{Z_{\beta_t}}{Z_{\beta_0}}.$$  

(14)

Here the work \(W\) is explicitly defined via Eq. (11) and

$$Q \equiv \int_0^t dt' \dot{x}_{\tau'} \frac{\partial}{\partial x_{\tau'}} H_S(x_{\tau'}, \tau').$$

(15)
can be regarded as a heat, which is transferred to the system. This definition of $Q$ can be understood based on the following consideration. The energy of the system can be changed by performing the work $W$ on the system or by supplying heat $Q$ to the system:

$$H_S(x_t, t) - H_S(x_0, 0) = \int_0^t dt' \frac{\partial}{\partial t'} H_S(x_{t'}, t') + \int_0^t dt' \frac{\partial}{\partial x_{t'}} H_S(x_{t'}, t').$$ (16)

Since the first term in this equation is the work (11), then the second term can be associated with the heat absorbed by the system.

Eq. (14) can be considered as the generalized Jarzynski equality. It means that (nonequilibrium) entropy can be changed by making some work and/or exchanging some heat. If we assume that the Hamiltonians $H$ and $H_S$ do not depend on time explicitly, then Eq. (14) reduces to

$$\langle \exp(- (\beta_S - \beta)Q) \rangle_{\rho_{\beta_0}} = 1,$$ (17)

which is the identity derived in Ref. [36]

**B. Systems in a steady state**

To derive the steady-state distribution, we can also apply the procedure of the external driving of the molecular Hamiltonian. [37, 38, 39, 40, 41, 42, 43, 44] Namely, we assume that the system-bath interaction is switched on adiabatically, so that the total time-dependent Hamiltonian reads as

$$H(\Gamma_t, t) = H_S(x_t) + H_B(X_t) + \exp(\varepsilon t)H_{SB}(x_t, X_t)$$ (18)

($\varepsilon$ is a positive infinitesimal). At a certain moment in the past $t = \tau \ll -1/\varepsilon$ our “system” and “bath” do not interact and are prepared at the temperature $T$ according to the grand canonical Gibbs distribution

$$\rho_{\tau} = Z_{\tau}^{-1} \exp(-\beta(H(\Gamma_{\tau}, \tau) - Y(\Gamma_{\tau}))).$$ (19)

Here

$$H(\Gamma_{\tau}, \tau) = H_S(x_\tau) + H_B(X_\tau), \ Y(\Gamma_{\tau}) = \mu_S N_S(x_\tau) + \mu_B N_B(X_\tau),$$ (20)

$\mu_\alpha$ are the chemical potentials for $S$ and $B$. $N_S(x_t) = 1$ if the spatial coordinate in the phase point $x_t$ belongs to the volume $V_S$ occupied by the system and zero otherwise; similarly for $N_B(X_t)$. 

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Preparation of the ensemble according to distribution (19) means that, initially, the system $S$ and the bath $B$ were in equilibrium with different heat baths. At the moment $\tau$, $S$ and $B$ are decoupled from their baths, and the $S - B$ interaction is gradually switching on. As a result of this incipient interaction, the systems can exchange their particles and energies with each other, so that the steady state is established at $t = 0$. To arrive at the desirable steady-state distribution, we can proceed as follows. First, we apply the thermodynamic limit to the bath degrees of freedom, so that $N_B \to \infty$, $V_B \to \infty$, $N_B/V_B \to \text{const}$ ($N_B$ is the number of bath particles and $V_B$ is the volumes occupied by the bath). Second, we tend $\varepsilon$ to zero (i.e., $\tau \to -\infty$) and propagate the initial equilibrium distribution (19) from $t = \tau$ to $t = 0$ with the time-dependent Hamiltonian (18). Third, we change from the phase variables $\Gamma$ (at $t = \tau = -\infty$) to $\Gamma_0 = \Gamma_0(\tau, t = 0)$. Then the non-equilibrium steady-state distribution at $t = 0$ can be written as follows:

$$\rho_{s0} = Z_{s0}^{-1} \exp(-\beta(H(\Gamma_0, 0) - Y(\Gamma_0))).$$

Here $Z_{s0}$ is the steady-state partition function and $Y(\Gamma_0)$ obeys the identity $\{Y(\Gamma_0), H(\Gamma_0, 0)\} = 0$. There are several equivalent forms of $Y(\Gamma_0)$. For the further discussion, the most elucidating is the following expression, which can be obtained if we substitute function $Y$ for $C$ in Eq.(9), set $t = 0$ and use the fact that the total number of particle is conserved ($d/dt(N_S + N_B) = 0$):

$$Y(\Gamma_0) = Y(\Gamma_\tau) + \Delta_\mu \mathcal{J}(\Gamma_0).$$

Here $\Delta_\mu = \mu_B - \mu_S$ and

$$\mathcal{J}(\Gamma_0) = \int_\tau^0 dt' \dot{\Gamma}_{t'} \frac{\partial}{\partial \Gamma_{t'}} Y(\Gamma_{t'})$$

is the time-integrated current density. The explicit form of $Y(\Gamma_0)$ makes evident a profound distinction between the equilibrium and steady-state preparation. If the chemical potentials of the system and the bath are the same, $\Delta_\mu = 0$, then the combined $S - B$ system will end up with an equilibrium distribution. Otherwise, the steady-state distribution establishes. It supports the steady-state currents, which are absent in equilibrium. It should be noted that Eq. (21) does not rely upon any sort of perturbation or linear response theory, and thus describes the steady-state distribution far from equilibrium. Furthermore, Eqs. (21)-(23) deliver an explicit formula for the steady-state distribution in terms of Hamiltonian (18).
We are in the position now to derive the work theorem for the steady-state systems. If we assume that $A = \rho_\tau$ and $B = \rho_{s0}$, we get then

$$\langle \exp(-\beta (W_{SB} - \Delta Y)) \rangle_{\rho_\tau} = \frac{Z_{s0}}{Z_\tau}. \tag{24}$$

Explicitly,

$$W_{SB} \equiv H(\Gamma_0, 0) - H(\Gamma_\tau, \tau) = \varepsilon \int_0^\tau dt' \exp(\varepsilon t') H_{SB}(x_{t'}, X_{t'}), \tag{25}$$

and

$$\Delta Y \equiv Y(\Gamma_0) - Y(\Gamma_\tau) = \Delta \mu J. \tag{26}$$

Eq. (24) allows us to follow the energy exchange during the onset of the steady state. By its definition (25), $W_{SB}$ looks similar to the nonequilibrium work in the standard Jarzynski equality (12). However, the quantity cannot be associated with the work performed on (by) the system. $W_{SB}$ can be regarded as the time-averaged value of the system-bath coupling $H_{SB}$. This follows immediately from the Abel’s theorem, \[37\] which states that the identity

$$\lim_{\varepsilon \to +0} \varepsilon \int_{-\infty}^0 dt' e^{\varepsilon t'} f(t') = \lim_{t \to \infty} \frac{1}{t} \int_{-t}^0 dt' f(t'), \tag{27}$$

holds for any function (operator) $f(t)$. On the other hand, $\Delta Y$ is proportional to the time-integrated current. Its presence in Eq. (24) is peculiar to the steady-state distribution, since $\Delta Y$ vanishes in equilibrium. The value of $\Delta Y$ equals the additional energy we have to spend for establishing the steady-state distribution.

An interesting result is obtained if we introduce the distribution

$$\rho_H = Z_H^{-1} \exp(-\beta (H(\Gamma_\tau, 0) - Y(\Gamma_\tau))), \tag{28}$$

$Z_H$ being the corresponding partition function. Eq. (28) assumes that the system and the bath are coupled all the time, but their chemical potentials are kept different. $\rho_H$ is neither an equilibrium nor the steady-state distribution. However, it is a perfectly legitimate mathematical object to consider. If we take $A = \rho_{s0}$ (21) and $B = \rho_H$ (28), we get

$$\langle \exp(-\beta \Delta \mu J(\Gamma_0)) \rangle_{\rho_{s0}} = \frac{Z_H}{Z_{s0}}. \tag{29}$$

Such a result can be obtained through the fluctuation theorems for currents,\[1, 4, 7, 52, 53, 54, 55\] and a similar formula has been derived in Ref. [41] in the context of the shear flow steady-state thermodynamics. The only difference between these results and ours is as
follows: \( Z_H/Z_{s0} \neq 1 \), in general. If the system-bath coupling is weak, however, we can write that \( Z_H/Z_{s0} = 1 + O(H_{SB}) \) and the ratio of the partition functions equals one in the leading order.

We assume now that our \( S+B \) system is prepared at \( t = 0 \) in the steady-state distribution \( \rho_{s0} \). Then, we switch the external field on at \( t = 0 \), so that the driven system Hamiltonian is explicitly described by Eq. (10) at \( t > 0 \). Under the influence of such a Hamiltonian, the steady-state distribution \( \rho_{st} \) will evolve into

\[
\rho_{st} = Z_{st}^{-1} \exp(-\beta(H(\Gamma_t, t) - Y(\Gamma_t))),
\]

\( t > 0 \). If we take \( A = \rho_{s0} \) (Eq. (21)) and \( B = \rho_{st} \) (Eq. (30)), we obtain

\[
\langle \exp(-\beta(W - (Y(\Gamma_t) - Y(\Gamma_0))) \rangle_{\rho_{s0}} = Z_{st}/Z_{s0}.
\]

Here the work \( W \) is explicitly defined via Eq. (11) and

\[
\Delta Y_s \equiv Y(\Gamma_t) - Y(\Gamma_0) = \int_0^t dt' \dot{\Gamma}_{t'} \frac{\partial}{\partial \Gamma_{t'}} Y(\Gamma_{t'}). \tag{32}
\]

Again, the additional term \( \Delta Y_s \) enters Eq. (32) as compared to its Jarzynski counterpart (12), manifesting thereby the necessity of additional energy expenses in the steady state. \( \Delta Y_s \) is always positive since it is the product of the current \( J \) and the chemical potential difference \( \Delta \mu \) and they always have the same sign.

C. Additional useful equalities and fluctuation theorems

(i). The (information) entropy associated with any nonequilibrium distribution can be defined as

\[
S_a \equiv -k_B \ln \rho_a. \tag{33}
\]

If we take \( A = \rho_a \) and \( B = \rho_b \) (the subscripts \( a \) and \( b \) correspond to any probability density function introduced above), then Eq. (11) tells us that

\[
\langle \exp((S_a - S_b)/k_B) \rangle_a = 1. \tag{34}
\]

This expression is thus very general and independent of a particular form of the nonequilibrium distribution. \[1, 4, 7\] It states that the path average of the exponential of the entropy production equals unity. For stochastic systems, a similar result has been proven in Refs.
It should be noted that, in general, \( S_a - S_b \) in Eq. (34) is the total system+bath entropy production, while the papers [27, 56] show that Eq. (34) holds for the entropy production of the system only, provided that the system dynamics is described by the Markovian master equation. It can be argued that if the bath is infinite then its entropy does not change and \( S_a - S_b \) is referred to the system. Furthermore, if \( A \) and \( B \) are distributions (5) or (13), then \( S_a - S_b \) is rigorously determined by the system operators only.

(ii). If we multiply Eq. (1) by \( \delta(\Gamma - \Gamma_t) \) and integrate over \( \Gamma_t \), we obtain the identity

\[
\int d\Gamma_0 A(\Gamma_0, 0) \delta(\Gamma - \Gamma_t) \left( A(\Gamma_0, 0)^{-1} B(\Gamma_t, t) \right) = B(\Gamma, t). \tag{35}
\]

If we take \( A \) and \( B \) to be the instantaneous Gibbs distributions (5), then we recover the expressions derived in Refs. [57, 58]. If \( A \) and \( B \) are certain nonequilibrium distributions, we arrive at the result derived in [59] for the system describing via an overdamped Langevin equation.

(iii). Let

\[
C(\Gamma_t, t) = \Psi(D(\Gamma_t, t)), \quad D(\Gamma_t, t) \equiv A(\Gamma_0, 0)^{-1} B(\Gamma_t, t), \tag{36}
\]

\( \Psi(D) \) being a certain function. We also introduce the inverse function, so that \( \Psi^{-1}(C) = D \). If we multiply Eq. (1) by \( \delta(w + C(\Gamma_t, t)) \) \((w \text{ being a parameter}) \), integrate it over \( \Gamma_t \), and use the notation (3), then we obtain the identity

\[
\Psi^{-1}(-w) \langle \delta(w + C(\Gamma_t, t)) \rangle_A = \langle \delta(w + C(\Gamma_t, t)) \rangle_B \tag{37}
\]

which can be coined as the generalized Crooks transient fluctuation theorem. If we take \( A \) and \( B \) to be the instantaneous Gibbs distributions (5), and let \( C \) be the ”minus” forward work performed on the system (11),

\[
C(\Gamma_t, t) = W(\Gamma_t, t) = -W(\Gamma_0, t) = H(\Gamma_0, 0) - H(\Gamma_t, t) = \frac{1}{\beta} \ln \left( D(\Gamma_t, t) \frac{Z_t}{Z_0} \right), \tag{38}
\]

then we obtain the Crooks transient fluctuation theorem [56, 57]

\[
\langle \delta(w - W(\Gamma_0, t)) \rangle_{\rho_0} = \langle \delta(w + W(\Gamma_t, t)) \rangle_{\rho_t} \exp(\beta w)Z_t/Z_0. \tag{39}
\]

Furthermore, let \( A \) and \( B \) be any (equilibrium or not) distributions evolving into each other along the forward and time-reversed trajectories, respectively. If we assume that \( C = \Psi(D) = k_B \ln D \) and adopt definition (33) for the entropy, then Eq. (37) yields the Crooks equality for the entropy production \( \Delta S(\Gamma_0, t) = -C(\Gamma_0, t) \):

\[
\langle \delta(w - \Delta S(\Gamma_0, t)) \rangle_{\rho_0} = \langle \delta(w + \Delta S(\Gamma_t, t)) \rangle_{\rho_t} \exp(w/k_B). \tag{40}
\]
Thus, as has been shown in Ref. [56], Eq. (40) is valid if we start from any, not necessary equilibrium, distribution. Eq. (40) has been derived in Ref. [33] for an externally driven Hamiltonian system coupled to several Hamiltonian heat reservoirs kept at different temperatures. This result is recovered if $A$ and $B$ are taken as nonequilibrium distributions (13).

(iv). We can generate complimentary work theorems by interchanging $A$ and $B$ in Eq. (2). For example, the so-obtained analogue of Eq. (29) reads

$$\langle \exp(\beta \Delta \mu J) \rangle_{\rho_H} = Z_s / Z_H.$$  

(41)

Therefore,

$$\langle \exp(\beta \Delta \mu J) \rangle_{\rho_H} \langle \exp(-\beta \Delta \mu J) \rangle_{\rho_{s0}} = 1.$$  

(42)

III. QUANTUM SYSTEMS

Almost all results obtained in the previous section are immediately transferable to the quantum case. In fact, we have to replace all the functions by operators in the Heisenberg representation (hereafter, the latter are marked by hats), Poisson brackets by commutators, and integrations by taking traces. Thus, our “generating expressions” (2) and (4) transform into

$$\text{Tr}(-i \hat{A}(0) \left[ \hat{A}(0)^{-1} \hat{B}(t) \right]) \equiv \langle \hat{A}(0)^{-1} \hat{B}(t) \rangle_A = \text{Tr} \hat{B}(t) \equiv B_t,$$

(43)

$$\langle \hat{A}(0)^{-1} \hat{B}(t) \rangle_A = B_t.$$  

(44)

Explicitly, the time dependence of any Heisenberg operator $\hat{C}$ is governed by the quantum analogue of the equation of motion (8)

$$\frac{d}{dt} \hat{C}(t) = \frac{\partial}{\partial t} \hat{C}(t) + i [\hat{H}(t), \hat{C}(t)].$$  

(45)

[...] is the commutator and we use the units with $\hbar = 1$. Thus

$$\hat{C}(t) - \hat{C}(\tau) \equiv \int_{\tau}^{t} dt' \hat{C}(t') = \int_{\tau}^{t} dt' \frac{\partial}{\partial t'} \hat{C}(t') + \int_{\tau}^{t} dt' i [\hat{C}(t'), \hat{H}(t')]$$

(46)
A. Systems at equilibrium

We write the total Hamiltonian as a sum of the system Hamiltonian, the bath Hamiltonian and their coupling and also allow for the system Hamiltonian to be explicitly time dependent,

$$\hat{H}(t) = \hat{H}_S(t) + \hat{H}_B + \hat{H}_{SB}. \quad (47)$$

If both $A$ and $B$ are the instantaneous quantum canonical distributions

$$\hat{A}(0) = \hat{\rho}_0 = Z_0^{-1} \exp(-\beta \hat{H}(0)), \quad \hat{B}(t) = \hat{\rho}_t = Z_t^{-1} \exp(-\beta \hat{H}(t)), \quad (48)$$

then we arrive at the expression

$$\left\langle \exp(\beta \hat{H}(0)) \exp(-\beta \hat{H}(t)) \right\rangle_{\rho_0} = Z_t/Z_0. \quad (49)$$

Now we can introduce the quantity

$$\hat{W} = \hat{H}(t) - \hat{H}(0) = \int_0^t dt' \frac{\partial}{\partial t'} \hat{H}(t') = \int_0^t dt' \frac{\partial}{\partial t'} \hat{H}_S(t'), \quad (50)$$

which is sometimes referred to as the work operator.\cite{14, 20, 21, 22} If $[\hat{H}(t), \hat{H}(0)] = 0$, then Eq. (49) transforms into the Jarzynski expression

$$\left\langle \exp(-\beta \hat{W}) \right\rangle_{\rho_0} = Z_t/Z_0. \quad (51)$$

If the Hamiltonians do not commute, then Eq. (49) can be rewritten as

$$\left\langle \exp(-\beta (\hat{W} + \hat{\delta} W)) \right\rangle_{\rho_0} = Z_t/Z_0, \quad (52)$$

where the quantum correction reads

$$\hat{\delta} W = -\frac{1}{\beta} \ln \left( \exp(\beta \hat{H}(0)) \exp(-\beta \hat{H}(t)) \right) + \hat{H}(0) - \hat{H}(t). \quad (53)$$

This quantum correction arises due to the fact that operators $\hat{H}(0)$ and $\hat{H}(t)$ do not commute in (49) and it is not associated with housekeeping heat.\cite{45, 46}

Alternatively, Eq. (49) can be recast into the form similar to Eq. (51) even for non-commutative $\hat{H}(0)$ and $\hat{H}(t)$, provided we introduce the chronological ordering operator $\mathcal{T}_<;\cite{21}$

$$\mathcal{T}_< \left\langle \exp(-\beta \hat{W}) \right\rangle_{\rho_0} = Z_t/Z_0. \quad (54)$$
Here $\hat{W}$, due to Eq. (50), is the work performed on the system $S$ only, in full analogy with the classical case. Eq. (54) makes it clear that the path average of the time-ordered Heisenberg operator $\exp(-\beta \hat{W}(t))$ yields the ratio of the partition functions. See also [1, 14, 15, 16, 17, 18, 19, 20, 21, 22] for the derivation of quantum Jarzynski equalities and discussion of various definitions of the quantum work operator.

If we consider the grand canonical ensemble, we arrive at a simple generalization of Eqs. (49) and (54) provided we assume that the chemical potential $\mu(t)$ is externally driven. We can take

$$\hat{A}(0) = \hat{\rho}_{N_0} = Z_{N_0}^{-1} \exp(-\beta(\hat{H}(0) - \mu(0)\hat{N})),$$
$$\hat{B}(t) = \hat{\rho}_{N_t} = Z_{N_t}^{-1} \exp(-\beta(\hat{H}(0) - \mu(t)\hat{N})).$$

(55)

$\hat{N}$ being the number of particles operator, $Z_{N_0}$ and $Z_{N_t}$ being the corresponding partition functions. Eq. (44) yields then

$$\langle \exp(-\beta(\hat{W} + \delta W - \Delta \mu(t)\hat{N})) \rangle_{\rho_{N_0}} = Z_{N_t}/Z_{N_0},$$

(56)

$\Delta \mu(t) = \mu(t) - \mu(0)$. If our Hamiltonian does not depend on time explicitly, then Eq. (56) simplifies to

$$\langle \exp(\beta \Delta \mu(t)\hat{N}) \rangle_{\rho_{N_0}} = Z_{N_t}/Z_{N_0},$$

(57)

irrespective of a particular driving protocol for $\Delta \mu(t)$.

Straightforward is to derive quantum analogues of other results obtained in Section IIA. For example, a quantum version of Eq. (14) reads

$$T_\beta < \langle \exp(-\beta S\hat{W} - (\beta S - \beta)\hat{Q}) \rangle_{\rho_{\beta_0}} > = Z_{\beta t}/Z_{\beta_0}.$$

(58)

Here the work operator is defined via Eq. (50) and the heat operator is determined as

$$\hat{Q} \equiv \int_0^t dt' i[\hat{H}(t'), \hat{H}_S(t')]$$

(59)

Note the chronological ordering in Eq. (58), which takes care of the fact that the operators do not commute, in general. We can also generalize Eq. (58) by changing from the canonical to grand canonical ensemble, thereby connecting the nonequilibrium work, the transferred heat, and the particle exchange to the ratio of two partition functions.
B. Systems in a steady state

Our consideration here parallels that of the classical systems in Sec. IIB. We assume that the system-bath interaction is switched on adiabatically, so that the total time-dependent Hamiltonian is written as

\[ \hat{H} = \hat{H}_S + \hat{H}_B + \exp(\varepsilon t)\hat{H}_{SB} \]  

(\( \varepsilon \) is a positive infinitesimal). At a certain time moment in the past \( t = \tau \ll -1/\varepsilon \) our “system” and “bath” do not interact and are prepared at the temperature \( T \) according to the grand canonical Gibbs distribution

\[ \hat{\rho}_\tau = Z_\tau^{-1} \exp(-\beta(\hat{H}(\tau) - \hat{Y}(\tau))) = Z_\tau^{-1} \exp(-\beta\hat{H}(\tau)) \exp(\beta\hat{Y}(\tau)). \]  

Here \( Z_\tau = \text{Tr}(\hat{\rho}_\tau) \),

\[ \hat{H}(\tau) = \hat{H}_S + \hat{H}_B, \quad \hat{Y}(\tau) = \mu_S\hat{N}_S + \mu_B\hat{N}_B, \]  

\( \mu_a \) are the chemical potentials for \( S \) and \( B \) and \( \hat{N}_a \) are the corresponding number operators. Apparently, \([\hat{H}(\tau), \hat{Y}(\tau)] = 0\).

If we apply the thermodynamic limit to the bath degrees of freedom (\( N_B \to \infty, V_B \to \infty, N_B/V_B \to \text{const} \)) and propagate the initial distribution \( \hat{\rho}_\tau \) from \( t = \tau \) to \( t = 0 \), we arrive at the (nonequilibrium) steady-state distribution at \( t = 0 \):

\[ \hat{\rho}_{s0} = Z_{s0}^{-1} \exp(-\beta(\hat{H}(0) - \hat{Y}(0))) = Z_{s0}^{-1} \exp(-\beta\hat{H}(0)) \exp(\beta\hat{Y}(0)). \]  

Here \( Z_{s0} = \text{Tr}(\hat{\rho}_{s0}) \) and, as in the classical case, \([\hat{H}(0), \hat{Y}(0)] = 0\). Explicitly,

\[ \hat{Y}(0) = \hat{Y}(\tau) + \Delta_\mu \hat{J}, \]  

where the time-integrated current operator \( \hat{J} \) is determined through the current operator \( \hat{J}(t) = \hat{N}_B(t) = i[\hat{H}(t), \hat{N}_B(t)] \) as

\[ \hat{J} = \int_\tau^0 dt' \hat{J}(t'). \]  

Distribution (63) can be derived within the framework of the method of statistical operator by Zubarev [37] and (generalized version of the) maximum entropy principle by Jaynes. [38] The use of distribution (63) and the standard Keldysh Non-equilibrium Green’s functions technique yields the same steady-state averages. [39, 42, 43] The equivalence between Keldysh non-equilibrium Green’s functions [60] and Zubarev method of statistical operator is discussed in the appendix.
Now we are in the position to derive quantum analogues of the expressions presented in Sec. 2B. If we assume that $\hat{A} = \hat{\rho}_\tau$ and $\hat{B} = \hat{\rho}_{s0}$, we obtain the expression

$$T_\zeta \left\langle \exp(-\beta(\hat{W}_{SB} - \Delta \hat{Y})) \right\rangle_{\hat{\rho}_\tau} = Z_{s0}/Z_{\tau}, \quad (66)$$

which imposes certain limits on the energy exchange during the onset of the steady state. Explicitly,

$$\hat{W}_{SB} \equiv \hat{H}(0) - \hat{H}(\tau) = \varepsilon \int_0^\tau dt' \exp(\varepsilon t') \hat{H}_{SB}(t'), \quad (67)$$

and

$$\Delta \hat{Y} \equiv \hat{Y}(0) - \hat{Y}(\tau) = \Delta \mu \int_\tau^0 dt' i[\hat{H}(t), \hat{N}_B(t)] = \Delta \mu \hat{J}. \quad (68)$$

As in the classical case, operator $\Delta \hat{Y}$ is responsible for the onset of the steady state, since $\Delta \mu \hat{J} = 0$ vanishes at equilibrium. A quantum counterpart of Eq. (29) reads

$$T_\zeta \left\langle \exp(-\beta \Delta \mu \hat{J}(\Gamma_0)) \right\rangle_{\rho_{s0}} = Z_H/Z_{s0}. \quad (69)$$

We can also write down a quantum extension of Eq. (32). Let our $S + B$ system be prepared at $t = 0$ according to the steady-state distribution (63). Then, we switch the external field on at $t = 0$, so that the driven system Hamiltonian is explicitly described by Eq. (47) at $t > 0$. Under the influence of this Hamiltonian, the steady-state distribution (21) will evolve into

$$\hat{\rho}_{st} = Z_{st}^{-1} \exp(-\beta(\hat{H}(t) - \hat{Y}(t))), \quad (70)$$

$t > 0$. If we take $\hat{A} = \hat{\rho}_{s0}$ (Eq. (63)) and $\hat{B} = \hat{\rho}_{st}$ (Eq. (70)), we get

$$T_\zeta \left\langle \exp(-\beta(\hat{W} - \Delta \hat{Y}_s)) \right\rangle_{\rho_{s0}} = Z_{st}/Z_{s0}. \quad (71)$$

Here the quantum work $\hat{W}$ is explicitly defined via Eq. (50) and

$$\Delta \hat{Y}_s = \hat{Y}(t) - \hat{Y}(0) = \int_0^t dt'[\hat{H}(t'), \hat{Y}(t')]. \quad (72)$$

If we define the quantum entropy operator as \[37, 38\]

$$\hat{S}_a \equiv -k_B \ln \hat{\rho}_a, \quad (73)$$

then the classical Eq. (34) remains valid in the slightly modified form:

$$\left\langle \exp(\hat{S}_a/k_B) \exp(-\hat{S}_b/k_B) \right\rangle_a = 1. \quad (74)$$
Here the subscripts $a$ and $b$ correspond to any density matrix operator introduced above. For a quantum system described by the Markovian master equation, a similar result has been derived in Ref. [17] for the entropy production of the system. In Eq. (74), $\hat{S}_a$ and $\hat{S}_b$ are the entropy operators for system+bath and the (strong) system-bath coupling can affect the system entropy production due to the quantum entanglement. By using the chronological ordering, Eq. (74) can be rewritten in the form similar to Eq. (54).

C. Experimental or computational verification of quantum identities

Within classical mechanics, it is conceptually straightforward to measure or compute the evolution of a certain physical quantity along the trajectory. Thus, putting aside technical and computational difficulties, verification and interpretation of the identities derived in Section II is primarily a matter of attributing a physical significance to the quantities $A$ and $B$. In quantum mechanics, it is not that clear how to measure or evaluate a certain physical quantity along the trajectory. An important question is therefore how the quantum identities can be interpreted, experimentally verified, or numerically tested for nontrivial systems. We discuss several possibilities in this section.

All our quantum identities can equivalently be recast into the so-called two-time measurement form. To this end, let us return back to our generating Eq. (43). Without any loss of generality, the operators $A$ and $B$ can be written in the exponential form

$$A(0) = \exp\{-\Lambda(0)\}, \quad B(t) = \exp\{-\Upsilon(t)\}, \quad (75)$$

here we use the Schrödinger representation for operators $A(0)$ and $B(t)$. We assume that the operators $\Lambda(0)$ and $\Upsilon(t)$ are Hermitian. Being the solutions of the eigenproblems for Hermitian operators

$$\Lambda(0) |\lambda\rangle = E_\lambda |\lambda\rangle, \quad \Upsilon(t) |v_t\rangle = E_{v_t} |v_t\rangle \quad (76)$$

($E_{v_t}$ and $|v_t\rangle$ depend on time parametrically) the eigenvectors are orthogonal and complete. We also assume that these eigenvectors span the same Hilbert space.

We define the evolution operator $G(0, t)$ in the Liouville space, which governs the time evolution of our system according to

$$G(0, t) |\lambda\rangle \langle \lambda| = |\lambda(t)\rangle \langle \lambda(t)|, \quad G^\dagger(0, t) B(t) \equiv G^\dagger(0, t) \exp\{-\Upsilon(t)\} \equiv \hat{B}(t). \quad (77)$$
\( G \) is unitary if the system dynamics is Hamiltonian, but it may be not if we consider the dissipative system dynamics. Then Eq. (43) can be rewritten in the following equivalent form:

\[
\left\langle \hat{A}(0)^{-1}\hat{B}(t) \right\rangle_A = A_0 \sum_{\lambda, \upsilon_t} \rho_{\lambda} |\langle \lambda(t) | \upsilon_t \rangle|^2 \exp\{-(E_{\upsilon_t} - E_{\lambda})\} \equiv A_0 \langle \exp\{-\Delta E\} \rangle = B_t. \tag{78}
\]

Here

\[
\rho_{\lambda} = A_0^{-1} \exp\{-E_{\lambda}\}, \quad A_0 = \text{Tr}\{A(0)\} = \sum_{\lambda} \exp\{-E_{\lambda}\}, \quad \Delta E \equiv E_{\upsilon_t} - E_{\lambda}. \tag{79}
\]

Eq. (78) can be interpreted in terms of the two-time measurement scheme. The first measurement at \( t = 0 \) selects an eigenfunction \( |\lambda\rangle \) of operator \( \Lambda(0) \) in the Schrödinger representation. The second measurement at time \( t \) selects an eigenfunction \( |\upsilon_t\rangle \) of operator \( \Upsilon(t) \) also in Schrödinger representation. The factor \( |\langle \lambda(t) | \upsilon_t \rangle|^2 \) gives us the probability of the system’s evolution from \( |\lambda\rangle \) to \( |\upsilon_t\rangle \). If we repeat the procedure many times, we yield the mean value of \( \langle \exp\{-\Delta E\} \rangle \), which is obtained by averaging over initial conditions and summation over final conditions. Thus, Eqs. (78) and (79) present an explicit measurement protocol, which (at least in principle) can be applied to test all our quantum equalities derived in Sections 3a and 3b. Of course, the meaning and interpretation of each equality depends on a particular form of (Hermitian) operators \( A \) and \( B \) (or \( \Lambda \) and \( \Upsilon \)), and a proper interpretation is not a trivial task. [62]

In several important cases, operators \( \Lambda \) and \( \Upsilon \) consist of sums of two or more operators. If the later operators commute (as, e.g., \( \hat{H} \) and \( \hat{N} \) in Eq. (55), \( \hat{H} \) and \( \hat{Y} \) in Eqs. (61) and (63)), then Eqs. (78) can be written in terms of the eigenvalues and eigenfunctions of the system Hamiltonian. This makes the interpretation of the equations much more physically transparent. Otherwise (as in Eq. (70)), the interpretation is less obvious.

In general, all our quantum work theorems are formulated through the (time-ordered) averages of Hermitian operators along the quantum trajectory. Thus, the corresponding expressions are well suited for the evaluation and/or testing by path integral numerical methods. Finally, the fluctuation theorems for steady-state currents and charge transport can directly be formulated in terms of observables, i.e., the probability density distributions of forward and backward currents. [7, 52, 53, 54, 55] This opens up a principal possibility of testing our Eq. (69), which can also be derived through the fluctuation theorems for currents.
IV. CONCLUSION

We have presented a unified and simple method for generating work theorems for classical and quantum Hamiltonian systems, both under equilibrium conditions and in a steady state. Throughout the paper, we adopt the partitioning of the total Hamiltonian into the system part, the bath part, and their coupling. We have rederived many equalities which are available in the literature and obtained a number of new results. All our expressions can be considered as rigorous mathematical identities, because the fulfillment of the Liouville theorem is required only.

The list of the work theorems is not exhausted by those presented in our paper, and new equalities can easily be generated via Eqs. (41) and (44), if necessary. Our results can be useful for obtaining various partition functions and (generalized) free energies through simulations and/or measurements performed on nonequilibrium systems. The (nonequilibrium) partition functions are important and useful quantities, since they can be employed exactly in the same manner as their equilibrium counterparts. For example, if we differentiate the logarithm of the steady-state partition function $Z_{s0}$ (which corresponds to either classical (21) or quantum (63) steady-state distribution) with respect to the difference of chemical potentials $\Delta \mu$, we get the steady-state value of the operator $\hat{Y}(0)$. This operator, which is linearly related to the time-integrated current $\hat{J}$, has the meaning of energy we have to spend for establishing the steady-state distribution.

Finally, we wish to comment on the role of nonconservative forces in establishing and maintaining steady states. Deriving our classical (21) and quantum (63) steady-state distributions, we did not invoke any external nonconservative forces for establishing the steady state. The only requirement is that the system under study (either with finite or infinite number of degrees of freedom) is coupled to the bath with infinite number of degrees of freedom, and the thermodynamic limit is applied. As a result, the system exchanges energy with the Hamiltonian bath during establishing and maintaining the steady state ($\Delta Y$ in Eqs. (21) and (63)) but no additional nonconservative dissipative forces are required. The distributions (21) and (63) can be derived within the framework of the method of statistical operator by Zubarev [37] and (generalized version of the) maximum entropy principle by Jaynes. The use of these distributions and the standard Keldysh Green’s functions technique yields the same steady-state averages. A controversial question is whether such an approach
is adequate for describing quantum transport on the nanoscale, but is beyond the scope of the present paper. This picture is in contrast with the steady-state thermodynamics in which non-conservative dissipative forces are responsible for establishing and maintaining the steady state, and thus an additional ”housekeeping heat” is necessary to keep the system in the steady state. So, beyond the formal general results, the equalities derived in the present article can be directly compared with their counterparts obtained within framework of the steady-state thermodynamics provided that the ”housekeeping heat” equals to zero. This simply renders the steady-state thermodynamics distributions equilibrium distributions. However, the non-conservative forces can straightforwardly be incorporated into our Hamiltonian approach if we switch to the thermostated dynamics.

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APPENDIX A: EQUIVALENCE OF ZUBAREV METHOD OF STATISTICAL OPERATOR AND KELDYSH NONEQUILIBRIUM GREEN’S FUNCTIONS

Within Zubarev method of statistical operator the steady state average of an operator $G$ is defined as

$$G = \text{Tr}[ho(0)G], \quad \text{(A1)}$$

where

$$\rho(0) = \lim_{\eta \to 0} \eta \int_{-\infty}^{0} dt' \exp(\eta t') U(t', 0) \rho_{\text{rel}}(t') U^\dagger(t', 0) \quad \text{(A2)}$$

Here $\rho_{\text{rel}}(t)$ is the ”relevant statistical distribution” and $U(t, 0)$ is the time evolution operator. [37] We assume that the relevant statistical distribution is given by the time-independent equilibrium density matrix

$$\rho_{\text{rel}}(t) = \rho(-\infty). \quad \text{(A3)}$$

We rewrite Eq.(A2) in the following form

$$\rho(0) = \lim_{\eta \to 0} \int_{-\infty}^{0} dt' \frac{d}{dt'}(\exp(\eta t')) U(t', 0) \rho(-\infty) U^\dagger(t', 0) \quad \text{(A4)}$$
Integrating Eq. (A4) by parts, we obtain

$$\rho(0) = U(-\infty, 0)\rho(-\infty)U^\dagger(-\infty, 0)$$  \hspace{1cm} (A5)

Therefore, the steady-state average value obtained within Zubarev method of statistical operator becomes

$$\mathcal{G} = \text{Tr}[U(-\infty, 0)\rho(-\infty)U^\dagger(-\infty, 0)G].$$  \hspace{1cm} (A6)

Since the averaging in Keldysh non-equilibrium Green’s functions is defined as

$$\text{Tr}[\rho(-\infty)U^\dagger(-\infty, 0)GU(-\infty, 0)],$$  \hspace{1cm} (A6)

it is clear that it can be obtained from Zubarev average (A6) by the cyclic permutation of the operators under the trace.

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For the steady state to occur, the system should normally be coupled to (at least) two different baths which are kept at different chemical potentials. To make the presentation simpler, we employ the (effective) single-bath description, since our formal expressions can be straightforwardly generalized to the multiple baths case.

If we consider $\rho_t$ for any finite $t$, then the classical and quantum evolution operators are unitary and thus $Z_\tau = Z_0$ for any finite system. Since we perform the thermodynamic limit $N_B \to \infty$, $V_B \to \infty$, $N_B/V_B \to \text{const}$ first and tend $\tau \to -\infty$ afterwards, then the evolution operators become isometric \cite{50} and, in general, $Z_\tau \neq Z_0$ (see Refs.\cite{37, 39, 42, 51} for more details).

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