Equilibration and temperature distribution in a driven ion chain

G-D Lin\textsuperscript{1,2,4} and L-M Duan\textsuperscript{1,3}

\textsuperscript{1}Department of Physics and MCTP, University of Michigan, Ann Arbor, MI 48109, USA
\textsuperscript{2}Department of Physics, University of Connecticut, Storrs, CT 06269, USA
\textsuperscript{3}Center for Quantum Information, IIIS, Tsinghua University, Beijing, People’s Republic of China
E-mail: lin@phys.uconn.edu

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Abstract. We study the non-equilibrium dynamics and equilibration in a dissipative quantum many-body system—a chain of ions with two points of the chain driven by a thermal bath under different temperatures. Instead of a simple linear temperature gradient (characterized by the local motional excitation) as one expects from a typical classical heat diffusion process, the temperature distribution in the ion chain shows surprisingly rich patterns, which depend on the rate of ion coupling to the bath, the location of driven ions and the dissipation rates of the other ions in the chain. By simulating the temperature evolution, we show that these unusual temperature distribution patterns in the ion chain can be quantitatively tested in experiments within a realistic time scale.

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4 Author to whom any correspondence should be addressed.
1. Introduction

Many-body non-equilibrium dynamics has attracted significant interest in recent years, in particular in connection with atomic experiments, where far-from-equilibrium phenomena can be conveniently investigated due to the long relaxation time of these systems [1, 2]. For instance, the dynamics after a quantum quench have been studied in several model Hamiltonians [3–6]. The physics becomes even richer if we add engineered dissipation to the underlying system, which is possible to realize in atomic experiments. Several interesting effects have been analyzed recently from the interplay between interaction and dissipation in cold atomic gases [7, 8].

Motivated by this line of research, in this paper we study how the system equilibrates (approaches a steady state) in a linear ion chain, with two points of the chain driven, respectively, by a heating and a cooling thermal bath. In a sense, the configuration here is an analogue of a classical example in thermodynamics—the heat propagation in a bar with its two ends fixed at different temperatures. To make a comparison, in our ion system we characterize the ‘local temperature’ for each individual ion with its average motional energy (the mean phonon number). This characterization is commonly used in quantum optics. We consider transverse phonon modes in this paper, and the transverse modes are only weakly coupled in an ion chain, so it makes perfect sense to introduce the concept of local temperature. In contrast with a simple linear temperature gradient as one sees in a classical bar, we find that the temperature distribution in a driven ion chain shows very rich behavior in its steady states: firstly, the temperature distribution depends critically on the ratio between the driving speed and the interaction rate in the ion chain. In the weak driving region, the temperature distribution is non-monotonic across the ion chain and shows a mirror effect. In the strong driving region, instead of a linear temperature distribution, all the ions between the two driven ones are stabilized to a constant temperature, which is in the middle of the two bath temperatures. Secondly, we find that the bath that drives the ion with a medium coupling rate plays a more effective role in determining the temperature of the other ions in the system. When the bath drives the ion strongly, it has little influence on the temperature of other ions in the chain, which is pretty counter-intuitive. To see the unusual phenomena predicted in this paper, we discuss the requirements for an experimental observation and calculate the time dynamics to achieve the steady-state temperature distribution. The qualitative features of the steady-state temperature distribution patterns are pretty insensitive to the size of the ion crystal and they show up already in a pretty small ion chain with fewer than ten ions. The relaxation time to the steady state is realistic for observation compared with the current experimental time scale.

This paper is organized as follows. In section 2, we provide the theoretical model and the calculation method. The main results on the temperature distribution patterns under different circumstances are presented and discussed in section 3. In section 4, we discuss the time dynamics to achieve the steady-state temperature distribution. Section 5 summarizes our major findings.

2. The model

We consider a chain of ions along the axial z-direction coupled with the Coulomb interaction and driven individually by a thermal bath. The thermal bath can be provided, for instance, by cooling or heating the laser beams shone on each ion. Both the driving rate and the effective bath
temperature can be tuned by controlling the intensity and the detuning of these laser beams. The oscillation of the \( i \)th ion around its equilibrium position is described by the coordinate and the momentum operators \( x_i, p_i \), with the dynamics determined by the Heisenberg–Langevin equation \([9, 10]\),

\[
\begin{align*}
\dot{x}_i &= p_i, \\
\dot{p}_i &= -\sum_j A_{ij} x_j - \gamma_i p_i + \sqrt{2\gamma_i} \zeta_i(t),
\end{align*}
\]

where \( A_{ij} \) denotes the coupling matrix between the ions, \( \gamma_i \) is the driving rate of the bath and \( \zeta_i(t) \) denotes the corresponding random force from the thermal bath. We consider in this paper the ions’ motion along the transverse \( x \) direction, so the coupling matrix \( A_{ij} \) is given by \( A_{ii} = \omega_i^2 - \sum_{j \neq i} 1/|z_j - z_i|^3 \) and \( A_{ij} = 1/|z_j - z_i|^3 \) for \( i \neq j \), where \( \omega_i \) is the transverse trapping frequency and \( z_i \) denotes the ion equilibrium position along the axial direction. We take \( \hbar^2/d_0 \) as the energy unit with the ion spacing \( d_0 \) and \( \omega_0 \triangleq \sqrt{\hbar^2/(md_0^2)} \) as the frequency unit so that every quantity in the matrix \( A_{ij} \) becomes dimensionless. Note that with the control of an anharmonic trapping potential along the axial direction, one can make the ion spacing uniform in a scalable trap \([11]\). For the conventional harmonic axial trap, the ion spacing is not uniform. In that case, the equilibrium positions \( z_i \) are determined numerically with the given trapping potential, and we take \( d_0 \) as the largest spacing in the ion chain. The temperature distribution pattern that will be shown in section 3 is insensitive to the details of the axial trapping potential, and for most of the calculations in the following, we assume that the ion spacing is uniform for simplicity (with exceptions in section 3 for the consideration of the time dynamics in a harmonic trap). For an independent Markovian bath, the random force \( \zeta_i(t) \) can be expressed as \( \zeta_i = -i\sqrt{\omega_i/2}(b_i - b_i^\dagger) \) with the bosonic field operator \( b_i(t) \) satisfying \( \langle b_i^\dagger(t_1)b_j(t_2) \rangle = T_\text{B}^i \delta_{ij} \delta(t_1 - t_2) \), where \( T_\text{B}^i \) is the average phonon number that characterizes the temperature of the bath, and \( \omega_i \triangleq \sqrt{A_{ii}} \) is the local oscillation frequency of the \( i \)th ion by fixing all other ions in their equilibrium positions. For typical linear ion traps with strong transverse confinement, we have \( \omega_i \approx \omega_x \ (\gg \omega_z, \text{the axial frequency}) \). The correlation of \( \zeta_i(t) \) is then given by \( \langle \zeta_i(t_1)\zeta_j(t_2) \rangle = \omega_i(T_\text{B}^i + 1/2)\delta_{ij} \delta(t_1 - t_2) \). In deriving this expression, we have used the relation \( \langle b_i^\dagger(t_1)b_j^\dagger(t_2) \rangle = 0 \) and \( \langle b_i(t_1)b_j(t_2) \rangle = 0 \) for a thermal bath.

The Langevin equation (1) can be solved exactly through diagonalization, with the solution formally expressed as

\[
q(t) = e^{-\Omega t} q(0) + \int_0^t d\tau e^{\Omega(t-\tau)} \eta(\tau),
\]

where

\[
q \triangleq (x_1, x_2, \ldots; p_1, p_2, \ldots)^T = \begin{bmatrix} \{x_i\} \\ \{p_i\} \end{bmatrix}, \quad \eta(t) \triangleq \begin{bmatrix} \{0\} \\ \{\sqrt{2\gamma_i} \zeta_i\} \end{bmatrix}
\]

and

\[
\Omega \triangleq \begin{bmatrix} 0 & -I \\ [A_{ij}] & [\gamma_i \delta_{ij}] \end{bmatrix}
\]
is a $2N \times 2N$ matrix, which can be diagonalized as $[U^{-1}\Omega U]_{\alpha\beta} = \lambda_{\alpha}\delta_{\alpha\beta}$. From this formal solution, we obtain the variance of the operators $x_i$ and $p_i$ as

$$\langle q^2 \rangle = \sum_{i=1}^{N} \sum_{\alpha,\beta=1}^{2N} U_{\mu\alpha} U_{\nu\beta} \left( e^{-(\lambda_{\alpha}+\lambda_{\beta})i} \left[ \langle x_i^2(0) \rangle U^{-1}_{\beta i} U^{-1}_{\alpha i} + \langle p_i^2(0) \rangle U^{-1}_{\beta,i+N} U^{-1}_{\alpha,i+N} \right] 
+ (1 - e^{-(\lambda_{\alpha}+\lambda_{\beta})i}) \frac{2\omega_i \gamma_i (T_i^B + \frac{1}{2})}{\lambda_{\alpha} + \lambda_{\beta}} U^{-1}_{\beta,i+N} U^{-1}_{\alpha,i+N} \right),$$

(3)

where $\mu = 1, 2, \ldots, N$ correspond to the $x$-operators and $\mu = N+1, N+2, \ldots, 2N$ correspond to the $p$-operators. The ‘temperature’ of each ion is similarly characterized by the average ‘phonon number’ of its local oscillation, so the temperature of the $i$th ion as a function of time $t$ is represented by $T_i(t) = \frac{1}{2} (\omega_i (x_i^2(t)) + \langle p_i^2(t) \rangle / \omega_i - 1)$. Here we have used $x_i = \frac{1}{\sqrt{2m}} (a_i + a_i^\dagger)$, $p_i = -i \sqrt{\frac{\hbar}{m}} (a_i - a_i^\dagger)$ and $T_i \triangleq \langle a_i^\dagger a_i \rangle$, where $x_i$ is in units of $\sqrt{\hbar/(m\omega_0)}$. As long as $\gamma_i$ is non-zero for some ions, each eigenvalue $\lambda_{\alpha}$ has a positive real part due to coupling of all the ions, and the system approaches a steady state as $t \rightarrow \infty$. The steady-state temperature for each ion is denoted by $T_i^s \triangleq T_i(t \rightarrow \infty)$.

3. Steady-state temperature distribution

In analogy to the example of heat propagation in a conducting bar, we consider a long ion chain with the two edge ions driven by different thermal baths with temperatures $T_1^B$ and $T_2^B$ (with $T_1^B < T_2^B$), respectively. For simplicity, we assume that the corresponding driving rate $\gamma_1 = \gamma_2 = \gamma$ and all ions in the middle experience no dissipation. After all the ions attain the steady state, the temperature distribution $T_i^s$ across the chain is shown in figure 1 for $N = 100$ ions under different driving rates $\gamma$. Apparently, the temperature distribution does not follow a linear gradient. There are three regions for the distribution of $T_i^s$, depending on the ratio between the driving rate $\gamma$ and the ion interaction rate. Note that in our units the interaction energy between the neighboring ions is of the order of unity. If $\gamma \gg 1$, the dissipation is much faster than the energy propagation in the chain, and without surprise the two edge ions have temperatures basically fixed by their corresponding bath temperatures $T_1^B$ and $T_2^B$. However, it is surprising that all the other ions in the middle approach almost the same temperature given by $T_i^s \simeq (T_1^B + T_2^B)/2$ for $i = 2, 3, \ldots, 99$ in this case. The temperature does not fall down gradually from the hot end to the cold end, as in the classical heat propagation problem, but takes a sharp jump right from the driven ion to the next one and then remains constant over the whole chain. In the opposite case of weak driving with $\gamma \ll 1$, the phonon propagation is faster than the bath driving, and all the ions approach the same temperature. It seems that the temperature distribution in this limit resembles a classical thermal equilibrium; however, this picture is not true. The example in figure 1 represents an exception instead of a rule where we put the cooling and heating ions exactly at the symmetric positions of an ion chain. As we will see in the following, when we shift the position of one of the ions to break the reflection symmetry, the temperature distribution in the weak driving limit has strange features. It is not even monotonic across the ion chain, in sharp contrast with the distribution from the diffusion process. Between these two limiting regions there lies a transition region with $0.01 < \gamma < 1$, where the temperature of the edge ions gradually approaches the corresponding
Figure 1. The steady-state ($t \to \infty$) temperature distribution $T^s_i$ (measured by the mean thermal phonon number) of a uniform ion chain with $N = 100$, where the 1st and the last (100th) ions are driven, respectively, by a cooling (at temperature $T_1^B = 2$) and a heating (at temperature $T_2^B = 10$) thermal bath under different driving rates $\gamma$. The temperatures of the driven ions (1st and 100th) are indicated by the thick curves. The other ions are assumed to be isolated from the bath, and the transverse trapping frequency is $\omega_x/\omega_0 = 10$.

bath temperature. Note that the temperature of the ions next to the driven ones follows a non-monotonic curve as one changes $\gamma$, although such a variation is pretty small.

To break the position symmetry of the cooling and the heating ions, we fix the cooling ion still at the edge, but move the heating ion inside the chain. Figure 2(a) shows an example of the temperature distribution where the heating ion is right at the center of the chain. When $\gamma \gg 1$, the heating ion separates two plateaus in the temperature distribution: a lower-temperature region between the heating and the cooling ions and a higher-temperature region on the other side of the heating ion. As $\gamma$ decreases, the two plateaus smoothly descend to the same temperature while the heating ion remains as a peak in the temperature distribution. When $\gamma \ll 1$, the temperature distribution has a reflection symmetry with respect to the chain center. A dip in temperature appears on the free edge of the chain (the left side of figure 2(a)), mirroring the cooling ion on the other side of the chain (the mirror effect). The temperature distribution is apparently non-monotonic in this case. The mirror effect exists for other positions of the heating ion. For instance, figure 2(b) shows the temperature distribution where the distance between the heating and the cooling ions is about one third of the chain length. Both the cooling and the heating ions have their mirror images in the temperature distribution in the weak driving limit. The mirror effect can be intuitively understood as follows. In the weak driving limit, the canonical modes for the ion motion are almost decoupled. The driving rate from the thermal bath on the edge ions can be decomposed into separate driving rates for each canonical mode, and each mode relaxes to its steady state. For an ion chain, the eigenfunction of each canonical mode in the real-space coordinate has a reflection symmetry with respect to the trap center. As a result, the temperature distribution for the final steady state also shows an (approximate) mirror symmetry in the weak driving limit.
Figure 2. The steady-state temperature distribution for $N = 101$ ions, where the cooling bath (at $T_1^B = 2$) remains at the first ion and the heating bath (at $T_2^B = 10$) moves to (a) the middle (51st) ion and (b) the 30th ion. The temperatures of the driven ions are indicated by thick (red) curves along the axis of the driving rate $\gamma$. The other parameters are the same as in figure 1. The temperatures profiles are highlighted by different thick (green) curves along the ion index axis to represent $\gamma = 10^{-3}, 10^{-2}, 0.1, 1$ and 10. For small $\gamma$, note that the temperature profiles in both (a) and (b) are found to have the mirror effect, i.e. a reflection symmetry of the distribution with respect to the middle (51st) ion.

In the above calculation, we assumed that $\gamma_1 = \gamma_2$ for the cooling and the heating ions. When these two driven ions are put at the edge, the middle ions are at a constant temperature $T_m$, which is exactly the average of the two bath temperatures. If the driving rate $\gamma_1 \neq \gamma_2$, we may expect that the bath that drives the ion more strongly plays a more important role in determining the temperature $T_m$ of the middle segment ions. This expectation, however, turns out to be not true. To show that, we map out the middle segment temperature $T_m$ in figure 3 as a function of $\gamma_1$ and $\gamma_2$. The figure shows that $\gamma_1 \approx 0.1$ is the optimal driving rate for which the corresponding bath has the largest influence on the temperature $T_m$. At this optimal value, the driving rate is comparable to the ion interaction rate in terms of the order of magnitude. The middle segment temperature is close to the temperature of the bath that drives the ion at the optimal rate. When the driving gets too strong or too weak, the bath plays little role in determining the temperature of the other ions in the chain. This result has important implications for the sympathetic cooling [12–14]: instead of fast cooling of the ancilla ions, cooling at a moderate optimal rate is more efficient in reducing the temperature of the computational ions.
So far we have neglected dissipation of the middle segment ions. Now, apart from the cooling and the heating bath (with temperatures $T_1^B$ and $T_2^B$, respectively) attached to the two edge ions, we assume that all the middle ions are coupled to a background bath with temperature $T_{bg}^B$ ($T_1^B < T_{bg}^B < T_2^B$) at a coupling rate $\gamma_{bg}$. The final temperature distribution of the ions is shown in figure 4 under different background coupling rates $\gamma_{bg}$. In this calculation, we fix $\gamma_1 = \gamma_2 = \gamma = 0.1$. For tiny $\gamma_{bg}$, the temperature distribution shows no noticeable difference compared with the case of $\gamma_{bg} = 0$. However, as $\gamma_{bg}$ increases to a moderate value with $\gamma_{bg}/\gamma \sim 0.01$, the temperature distribution of the middle segment ions has a clear linear spatial gradient, resembling the linear temperature distribution of a classical bar in the heat diffusion problem. As $\gamma_{bg}$ further increases and gets close to $\gamma$, the temperature of the middle ions is pinned to the background bath temperatures $T_{bg}^B$ as one expects and the temperature gradient disappears again in this limit.

4. Temperature evolution

We note that the temperature distributions shown in this paper are insensitive to the size of the ion system. We have checked the temperature distribution of the ions with the size of the chain varying from ten to a few hundreds of ions and noted no qualitative change in the distribution pattern. To experimentally test these unusual temperature distribution patterns, one can start with a small system of a few ions that are within the reach of the current experimental technology. The final temperature (the mean phonon number of the ion motion) can be detected, for instance, by measuring the scattering sidebands of a laser beam through an ion. The asymmetry in the blue and the red sidebands and their ratio give direct inference of the thermal phonon number of the ion motion \[15–17\].
To probe the properties of the steady states, we need to know the time scale to approach these steady states. Figure 5 shows the relaxation dynamics for 20 ions in either a harmonic or an anharmonic uniform trap [11]. For a uniform trap (in figure 5(a)), the calculation shows that there are two time scales in the equilibration. First, with a time scale $t_1 \sim 1/\gamma$, the temperature of the two edge ions quickly approaches the corresponding bath temperature. During this step, the temperature of the middle segment ions only changes slightly, and the change gets smaller as one moves away from the driven ions. After that, a longer time scale $t_2$ sets in, representing the interaction-driven equilibration process. All the ions gradually approach the steady-state temperature. Note that the temperature of the edge ions (as well as the other ions that are close to the two driven ones) does not follow a monotonic evolution curve. Instead, it first comes pretty close to the corresponding bath temperature and then is dragged back toward its steady-state value. The value of the second time scale $t_2$ increases with the system size (roughly linearly) and $t_2 \sim 40/\gamma$ for 20 ions. For an ytterbium ion ($^{171}$Yb$^+$) chain with spacing 10 $\mu$m, $t_2$ is about 3 ms, which is a pretty reasonable time scale for experiments. For a harmonic trap (in figure 5(b)), the basic feature is similar except that the second time scale $t_2$ becomes site dependent.
Figure 5. The temperature relaxation dynamics for an ion chain with $N = 20$ in (a) a uniform anharmonic trap, (b) a harmonic non-uniform trap and (c) a harmonic trap with a background coupling rate $\gamma_{bg}/\gamma = 10^{-3}$ and a background temperature $T_{B}^{B} = 4$. The 1st and the 20th ions are driven, respectively, by a cooling (at $T_{B}^{B} = 2$) and a heating (at $T_{B}^{B} = 10$) bath, with the same driving rate $\gamma_{1} = \gamma_{2} = \gamma = 0.1$, as indicated by the thick (red) curves. The initial temperature is assumed to be $T_{i}(t = 0) = 5$ for all ions. The transverse frequency $\omega_{x} = 10$ in units of $\sqrt{e^{2}/(md_{0}^{3})}$, where $d_{0}$ is the largest spacing in the ion chain (between the 1st and 2nd ions or between the 19th and 20th ions) for the harmonic cases. In terms of real numbers, with a typical choice of $d_{0} = 10 \mu m$ for 20 ytterbium ions in a harmonic trap, the trapping frequencies for the transverse and the axial traps are given, respectively, by $\omega_{x} \approx 2\pi \times 1.4 \text{ MHz}$ and $\omega_{z} \approx 2\pi \times 76.5 \text{ kHz}$, and the driving rate $\gamma \approx 14 \text{ kHz}$.

The edge ions and their neighbors approach the corresponding steady-state temperature with a time scale that is comparable to the case of a uniform chain. However, as one moves away from the edge, the equilibration time gets much greater with an exponential increase. For the middle ion, the equilibration is not finished yet with $t \sim 10^{10}/\gamma$. With such an extremely long time scale, of course, one cannot neglect the small dissipation of the other ions to the background bath. If we take into account a small background dissipation, e.g. with a rate $\gamma_{bg}/\gamma \sim 10^{-3}$
as shown in figure 5(c), the long tail in the temperature evolution is completely gone for the harmonic trap. Now, the temperature of all the ions approaches their steady values within a time scale that is comparable to the uniform case, although the steady-state temperature of the middle segment ions is dragged down a bit toward the background bath temperature.

5. Conclusion

In summary, we have shown that the steady-state temperature distribution of a driven ion chain shows surprisingly rich patterns. Many of the features of these patterns are unexpected, and are in sharp contrast with the simple linear temperature gradient as one sees in the classical heat diffusion problem. Our calculation is based on exact numerical methods without any unreliable approximations, so we believe that all the unusual temperature distribution patterns revealed by this calculation will show up in experiments, although we still lack an intuitive explanation of some of these features. We also investigate the relaxation dynamics and the time scale to reach the steady state, and show that these patterns should be observable within a realistic time scale in a small system that is within the current experimental reach.

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