Mutual information in coupled double quantum dots

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Abstract. Coupled bistate multipartite systems are extensively studied as a versatile model in the field of information thermodynamics. As such systems are functionalized via the interaction among the particles, mutual information is a key property in their operation. In this study, we analyze a simple numerical model comprising a set of coupled double quantum dots, where the disconnection of the elements is represented by the removal of Coulomb interaction between the quantum dots. We calculate the mutual information in the model system, as the Kullback–Leibler divergence between the connected and disconnected status, through the probability distribution of the electronic states from the master transition-rate equations. We reasonably demonstrate that the increase in the strength of interaction between the quantum dots leads to higher mutual information, owing to the larger divergence in the probability distributions of the electronic states. Our numerical model could be a useful basic tool for the investigations of information-driven systems.

Keywords: numerical simulations, quantum dots, stochastic thermodynamics, systems neuroscience
1. Introduction

In the context of information thermodynamics [1–5], the mutual information is defined as the difference between the actual Shannon entropy of the system and that in the absence of the correlations among the random variables, and represents the intensity of inter-variable correlations [6–10]. In other words, mutual information is the divergence between the actual probability distribution and that for the independence of the variables, to quantify the reduction in uncertainty for the states of the system on making measurements. Bipartite, four-state configurations are a handy model employed for the investigations in the field of information thermodynamics [11–16]. Quantum dots [17–21], often referred to as artificial atoms, are an adopted candidate for a material component of such setups [11, 22–27]. The discretized density of states in quantum dots, owing to the three-dimensional confinement of electrical carriers, enables single-electron manipulation [28–30], which may make the discussions clear in information statistical mechanics. We previously studied a coupled double quantum dot system as an autonomous information engine, calculating the steady-state entropy production rate in each component, heat and electron transfer rates via the probability distribution of the four electronic states from the master transition-rate equations, to acquire device-design principles toward the realization of corresponding practical energy converters [31]. As this system strongly relies on the interaction between the quantum dots, mutual information is a key property in the operation of information engines. In the present study, we carry out simulations to analyze the mutual information between the quantum dots, to provide a simple but practically useful model setup for numerical investigations of information-driven systems.

2. Theory and calculation methods

The model setup of the present study comprises two quantum dots and three thermal/electronic reservoirs around as schematically depicted in figure 1. Each quantum dot can contain up to one electron. One quantum dot with an electronic potential energy
\( \varepsilon_X \) functions as an electronic detector by ‘checking’ whether an electron is in the other quantum dot through capacitive interaction strength or Coulomb interaction energy \( U \) between the two quantum dots. This ‘detector dot’ is kept at a temperature \( T_D \) and connected to thermal and electronic reservoirs, both having the same temperature \( T_D \) and an electronic potential energy \( \mu_D \). The other quantum dot with an electronic potential energy \( \varepsilon_Y \) is connected to two reservoirs through electrical leads and enables an electrical current flow. This ‘system dot’ is kept at a temperature \( T_S \) and connected to two thermal and electronic reservoirs both at \( T_S \) with electronic potential energies \( \mu_H \) and \( \mu_L \) (\( \mu_H > \mu_L \)). It should be noted that by such a connecting configuration, this quantum-dot system realizes a non-equilibrium steady state, and is productively active, performing work in exchange for the decrease in thermal energy (the thermoelectric engine mode operation) or the increase in entropy (the environmental information engine mode operation) of the reservoirs, as living bodies [31]. The potential-energy relations among the components in the setup are schematically shown in figure 2 for clarification. This double-quantum-dot configuration as a whole can drive electrical current between the reservoirs of \( \mu_H \) and \( \mu_L \) through the system dot, even in the direction from \( \mu_L \) to \( \mu_H \) against the potential slope and thus generate work by properly setting the transition or tunneling rates across the interfaces between the quantum dots and reservoirs, as shown in reference [31]. Each quantum dot has an electronic state 0 or 1, where 1 and 0 mean that the dot is filled or not filled (i.e., empty) with an electron, respectively. In this way, the total electronic state \((x, y)\) will be \((0, 0)\), \((0, 1)\), \((1, 0)\), or \((1, 1)\). For the state \((1, 1)\), the electronic potential energy in the quantum dots will increase to \( \varepsilon_X + U \) and \( \varepsilon_Y + U \) for the detector and system dots, respectively, due to Coulomb repulsion. We set the time resolution fine enough so that no simultaneous or diagonal jump, such as a transition from \((0, 0)\) to \((1, 1)\), is assumed in our bipartite setup.
The time evolution of the probability of the state \( p(x, y) \) can be generally written as a master equation:

\[
d_t p(x, y) = \sum_{x', y'} \left\{ \frac{W_{x, y'}^{y', y} p(x', y') - W_{x', y'}^{x', y} p(x, y)}{d_t} \right\}, \tag{1}
\]

where \( W_{x, y'}^{y', y} \) is the transition rate from a state \((x', y')\) to \((x, y)\) and we have:

\[
W_{10}^y = \Gamma f_y \tag{2}
\]

and

\[
W_{01}^y = \Gamma (1 - f_y) \tag{3}
\]

for the electron transfer on the detector dot and

\[
W_{x, y'}^x = \Gamma_x f_x^y \tag{4}
\]

and

\[
W_{x, y'}^{0, x} = \Gamma_x f_x^{x'} \tag{5}
\]

for the system dot in this model. Note again that for the jumps, either \( x \) or \( y \) is fixed at each time step. \( \Gamma \) is the electronic tunneling rate between the detector dot and its reservoir. We assume the density of states in the detector-side reservoir to be uniform so that \( \Gamma \) is independent of \( y \). \( \Gamma_x^y \) is the tunneling rate between the system dot and its reservoirs where \( v = H \) or \( L \) corresponds to the higher- or lower-potential reservoir, respectively. In contrast, we assume nonuniform profiles of the density of states in the system-side reservoirs so that \( \Gamma_x^v \) depends on \( x \). Fermi distribution functions for the detector and system dots have forms of:

\[
f_y = \frac{1}{1 + \exp \left( \frac{\varepsilon + y U - \mu_D}{T_D} \right)} \tag{6}
\]

and

\[
f_x^v = \frac{1}{1 + \exp \left( \frac{\varepsilon + x U - \mu_v}{T_s} \right)} \tag{7}
\]

respectively. For simplicity, the Boltzmann constant is set to unity or absorbed into the temperatures throughout this paper. We then determine the steady-state probability.
distribution of \( p(x, y) \) for the four electronic states through the master transition-rate equations of equation (1).

For the disconnected model, we consider the probability distribution of the states \( q(x, y) \) in the system under consideration that corresponds to \( p(x, y) \) for the case \( U = 0 \), where the interactive information between the quantum dots is removed. Incidentally, as a consequence of \( U = 0 \), \( q(x, y) = q(x)q(y) \) holds, where \( q(x) = q(x, 0) + q(x, 1) \), \( q(y) = q(0, y) + q(1, y) \). We then define the mutual information in the coupled double quantum dot system, via the Kullback–Leibler divergence, as:

\[
\Phi_{MI} = D_{KL} \{ p(x, y) \| q(x, y) \} = \sum_{x,y} p(x,y) \ln \frac{p(x,y)}{q(x,y)}
\]

\[
= p(0,0) \ln \frac{p(0,0)}{q(0,0)} + p(0,1) \ln \frac{p(0,1)}{q(0,1)} + p(1,0) \ln \frac{p(1,0)}{q(1,0)} + p(1,1) \ln \frac{p(1,1)}{q(1,1)}
\]

where \( D_{KL} \{ p(x, y) \| q(x, y) \} \) is the Kullback–Leibler divergence between the probability distributions \( p(x, y) \) and \( q(x, y) \).

### 3. Results and discussion

Figure 3 presents a set of our calculation results for the mutual information of the coupled double quantum dot system in dependence on the capacitive interaction strength or quantum-dot Coulomb repulsion energy \( U \) for varied temperatures of the detector side \( T_D \), for the cases \( \mu_D = \epsilon_X - U/2 \) and \( \mu_D = \epsilon_X + U/2 \) (adopted from reference [31]). It should be noted that, because there are too many conditional parameters to change for the numerical calculations, we intentionally provided some restrictions among the parameters in this study. It is observed for all \( T_D \)'s that as \( U \) increases, the mutual information increases. This resulted trend seems reasonable in the view of the integration of information in the system; stronger interaction between the quantum dots provides a higher degree of integrated information. The mutual information is observed to be strongly dependent on \( T_D \) or the relative difference between \( T_S \) and \( T_D \). Lower \( T_D \) provides higher mutual information for a given \( U \). This trend can be attributed to the fact that under the fixed-\( T_S \) conditions, lower \( T_D \) and thus larger difference between \( T_S \) and \( T_D \) provides higher electron-transport selectivity, functioning towards the same directionality as larger \( U \). With no interaction between the quantum dots, \( U = 0 \), the four probabilities of the states almost equivalently share the pie, \( q(x, y) = 0.25 \). In the large-\( U \) regime, the repulsive interaction between the quantum dots tends to eliminate \( p(1,1) \). For the condition \( \mu_D = \epsilon_X - U/2 \), for larger \( U \), \( p(0,0) \) and \( p(0,1) \) approach 0.5 while the other probabilities of the states go to zero, owing to the deep reservoir level for the detector dot and \( \epsilon_Y \) locating in the middle of \( \mu_H \) and \( \mu_L \) for the system dot. This situation provides the limit of mutual information as \( 2 \times 0.5 \times \ln(0.5/0.25) = \ln 2 - 0.693 \), as observed in figure 3(a). On the other hand, for the condition \( \mu_D = \epsilon_X + U/2 \), for larger \( U \), \( p(1,0) \) approaches unity while the others go to zero, because of the...

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Mutual information $\Phi_{MI}$ in the coupled double quantum dot system in dependence on the capacitive interaction strength between the quantum dots $U$ for various temperatures of the detector quantum dot $T_D$ for the cases (a) $\mu_D = \varepsilon_X - U/2$ and (b) $\mu_D = \varepsilon_X + U/2$ under the condition $\varepsilon_X = \varepsilon_Y = 1$, $\mu_H = 1.1$, $\mu_L = 0.9$, $T_S = 1$, $\Gamma = 100$, $\Gamma_0^H = \Gamma_1^L = 10$, and $\Gamma_1^H = \Gamma_0^L = 0.1$. QD stands for quantum dot.

Figure 4. Mutual information $\Phi_{MI}$ in dependence on the capacitive interaction strength between the quantum dots $U$ for various tunneling rates $\Gamma_H^0$ and $\Gamma_L^1$, under the condition $\varepsilon_X = \varepsilon_Y = 1$, $\mu_H = 1.1$, $\mu_L = 0.9$, $T_D = 0.1$, $T_S = 1$, $\Gamma = 100$, and $\Gamma_1^H = \Gamma_0^L = 0.1$.

We plot the mutual information for various tunneling rates in figure 4. We varied the tunneling rates for the electron-flow direction against the potential differences $\Gamma_0^H$ and $\Gamma_1^L$ while fixing the tunneling rates in the direction down the potential slopes $\Gamma_1^H$ and $\Gamma_0^L$.

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Figure 5. Mutual information $\Phi_{\text{MI}}$ in dependence on the capacitive interaction strength between the quantum dots $U$ for various relative energy-level positions $\mu_H$ and $\mu_L$ to $\varepsilon_Y$, keeping $\mu_H - \mu_L = 0.2$ constant, under the condition $\varepsilon_X = \varepsilon_Y = 1$, $T_D = 0.1$, $T_S = 1$, $\Gamma = 100$, $\Gamma_0^H = \Gamma_1^L = 10$, and $\Gamma_1^H = \Gamma_0^L = 0.1$.

As seen in this set of mutual-information results, the mutual information is moderately dependent on the antisymmetric relative amplitudes of the electronic tunneling rates, i.e., that $\Gamma_0^H$ and $\Gamma_1^L$ are larger than $\Gamma_1^H$ and $\Gamma_0^L$. This type of antisymmetry was essential for the operation of the coupled double quantum dot system as an information engine and strongly influenced the information-engine efficiency [31]. However, it is observed that this is not the case for the mutual information, and it works even for the nonsymmetric case, $\Gamma_0^H = \Gamma_0^L = \Gamma_1^H = \Gamma_1^L = 0.1$.

Here, we investigate the influence of the energy levels’ relative positions among the system dot and its reservoirs. Figure 5 shows the mutual information of the coupled double quantum dot system with varied $\mu_H$ and $\mu_L$ to $\varepsilon_Y$, keeping $\mu_H - \mu_L$ constant. The mutual information is observed to be significantly influenced by the relative positions of the energy levels of $\mu_H$ and $\mu_L$ to $\varepsilon_Y$. For all of the three cases, $p(1, 0)$ approaches to unity for larger $U$, which is the dominant factor for the mutual information. On the other hand, $q(1, 0)$ is smaller for higher $\mu_H$ and $\mu_L$ relative to $\varepsilon_Y$, owing to the dominance by $q(0, 1)$ and $q(1, 1)$, which provides higher mutual information through equation (8). The plateau in the region $U = 1–2$ observed in the curve for the condition $\mu_H = 3.1$, $\mu_L = 2.9$ can be explained as follows. Figure 6 plots $p(x, y)$ in dependence on $U$ for the condition $\mu_H = 3.1$, $\mu_L = 2.9$. As observed, the evolution of $p(0, 1)$, which is the dominant factor for the increase in the mutual information in the small-$U$ regime, by $U$ is damped around the region $U = 1–2$ because the electronic energy level $\varepsilon_Y + U$ approaches $\mu_H$ and $\mu_L$. This situation of the relation of energy levels starts the escape of the electron from the system dot out to its reservoirs, and allows the entrance of the electron into the detector dot from its higher reservoir level, in the existence of the electronic inter-dot Coulomb repulsion by $U$. By contrast, the conditions $\mu_H = 0.2$, $\mu_L = 0.4$ and $\mu_H = 1.1$, $\mu_L = 0.9$ exhibit no such an intermediate plateau in the evolution of mutual information by $U$, as observed in figures 3–5, because $\varepsilon_Y + U$ is always higher than $\mu_H$ and $\mu_L$. After the point $U = 2$, where $\varepsilon_Y + U$ overlaps with $\mu_H$ and...
$\mu_L$, $p(1, 0)$ steeply increases and becomes the dominant factor to recover the increase in the mutual information until reaching unity. Incidentally, it is observed in figure 6 that after the conversion of the gradients between $p(0, 1)$ and $p(1, 0)$ around $U = 1-2$, as explained above, $p(1, 0)$ overtakes $p(0, 1)$ exactly at the point $U = 4$, where the electronic potential energy difference $\mu_D - \varepsilon_X$ for the detector dot equals (the middle of $\mu_H$ and $\mu_L$) $- \varepsilon_Y$ for the system dot, and thus the absorbing driving forces into the dots are leveled.

4. Conclusions

In this study, we carried out a series of numerical simulations for the mutual information in a model coupled double quantum dot system with an artificial function to ‘check’ the electronic state of each other via the Coulomb interaction. The evolutions of the mutual information of the system in dependence on $U$ for various setup conditions were quantitatively analyzed in relation to the probability distributions of the electronic states. It was reasonably observed that the mutual information increases with $U$, because stronger interaction between the quantum dots provides a higher degree of integrated information. The mutual information of the system was observed strongly dependent on the relative temperatures of the detector and system sides owing to the influence of the electron-transport selectivity, while moderately insensitive to the antisymmetric relative amplitudes of the electronic tunneling rates. The mutual information is also found to be significantly influenced by the relative positions of the energy levels of $\mu_H$ and $\mu_L$ to $\varepsilon_Y$, due to the drastic turnover of the dominant electronic states. Our numerical model could be a simple but useful numerical tool for the future investigations of information-driven systems.
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