Abnormal energy metabolism can alter foraging behavior in termites in different social contexts

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Abstract

Foraging behavior, as an energy-consuming behavior, is very important for collective survival in termites. How energy metabolism related to glucose decomposition and ATP production influences foraging behavior in termites is still unclear. Here, we analyzed the change in energy metabolism in the whole organism and brain after silencing the key metabolic gene isocitrate dehydrogenase (IDH) and then investigated its impact on foraging behavior in the subterranean termite Odontotermes formosanus in different social contexts. The IDH gene exhibited higher expression in the abdomen and head of O. formosanus. The knockdown of IDH resulted in metabolic disorders in the whole organism, including the impairment of the NAD\(^+\)-IDH reaction and decreased ATP levels and glucose accumulation. The dsIDH-injected workers showed significantly reduced walking activity but increased foraging success. Interestingly, IDH downregulation altered brain energy metabolism, resulting in a decline in ATP levels and an increase in IDH activity. Additionally, the social context obviously affected brain energy metabolism and, thus, altered foraging behavior in O. formosanus. We found that the presence of predator ants increased the negative influence on the foraging behavior of dsIDH-injected workers, including a decrease in foraging success. However, an increase in the number of nestmate soldiers could provide social buffering to relieve the adverse effect of predator ants on worker foraging behavior. Our orthogonal experiments further verified that the role of the IDH gene as an inherent factor was dominant in manipulating termite foraging behavior compared with external
social contexts, suggesting that energy metabolism, especially brain energy metabolism, plays a crucial role in regulating termite foraging behavior.

**Author summary**

Foraging behavior plays a key role in collective survival in social insects, as found in termites. Worker termites are responsible for foraging duty and exhibit large foraging areas and long foraging distances, so they need to consume much energy during foraging. It is well established that energy can influence insect behaviors. However, how energy metabolism affects foraging behavior in termites remains unknown. Here, we found that the downregulation of the conserved metabolic gene *IDH* impaired whole-organism and the brain energy metabolism and further altered foraging behavior, resulting in decreased walking activity but increased foraging success in the termite *O. formosanus*, which is an important insect pest damaging embankments and trees in China. Additionally, the social context affected brain energy metabolism and obviously changed foraging behavior in *O. formosanus*, causing a decline in foraging success in the absence of nestmate soldiers and the presence of predator ants. However, the increasing number of nestmate soldiers strengthened social buffering to relieve the negative effect of predator ants on worker foraging behavior. Our findings provide new insights into the underlying molecular mechanism involved in modulating the sophisticated foraging strategy of termites in different social contexts from the perspective of energy metabolism.
Introduction

All behaviors consume energy (to various degrees), and some also facilitate energy acquisition, such as foraging behavior, which presents a cost-benefit trade-off between growth and survival in organisms [1, 2]. The foraging behavior of social insects is an extremely complex process involving the self-organization of a large number of individuals to collect foods from various sources [3, 4]. For example, the honey bee *Apis mellifera* recruits additional foragers to valuable resources by dancing to signal the location of the resources to their nestmates [5]. In seed-harvesting ants of *Pogonomyrmex* spp., a mixture of individual and group foraging is the most common foraging strategy [6]. A group of foraging workers utilizes clay to build structural supports to increase foraging resources in the termite *Coptotermes acinaciformis* [7]. Therefore, social insects prefer to employ highly effective and energy-saving foraging strategies to obtain sufficient food and thereby fulfil the energy requirements of their colonies.

A large body of research is focused on the relationship between energy metabolism and behavior in both vertebrates and invertebrates [1, 8-10], revealing that the brain is a major regulator of behavior and metabolic processes [11, 12]. For instance, in adult *Drosophila*, the knockdown of glycolytic enzymes specifically in the glial cells of the nervous system in the brain leads to severe locomotor deficits [13]. The regulation of behavior is affected by the nerves and requires large amounts of energy in the brains of
the cricket *Gryllus campestris* L. and the ant *Pheidole rhea* [14, 15]. It seems that the basic energy metabolism related to neuromodulation can explain the relationship between energy metabolism and behavior [16, 17]. However, the role of energy metabolism in regulating social behaviors is still unclear.

Complex social contexts can affect foraging behavior in social insects. Considerable research has shown that predation risk seriously affects animal activity and foraging behavior [18-22]. Biological individuals may make collective decisions in intricate social contexts [23] and then respond to the presence of predators by showing a range of behavioral and physiological changes to reduce the likelihood of injury or death [24]. For instance, cryptic termites can change their foraging strategy by eavesdropping on vibrational cues from the footsteps of predatory ants to limit predation risk [25, 26]. Honey bees avoid flowers with crab spiders and flowers that have recently held spiders during foraging [27]. Therefore, predation pressure from predators is an important factor impacting foraging behavior in social insects. In addition, social information provided by nestmates can also change individual foraging decisions [28]. In social insects, soldiers play a vital role in coping with the risk of external predation by sending out warning signals in a special way so that the other nestmates can avoid risks in a limited time through nestmate recognition during the process of foraging [29-31]. Social buffering is involved in the ability of neighboring individuals in a colony to reduce the negative impact of stressors on other individuals in a wide range of vertebrates [32]. For example, in the termite *Reticulitermes flavipes*, the presence of
nestmate soldiers can change the ability of workers to cope with the competition risk imposed by conspecific non-nestmates [24]. However, the changes in behavioral characteristics are unclear when nestmate soldiers provide social buffering to foraging termites with different energy levels in the presence of predator ants.

Theoretically, both inherent genes and external social contexts can influence foraging behavior in insects. The foraging gene (for), encoding a cGMP-dependent protein kinase (PKG), is related to the foraging behavior of several social and solitary insects, including the lower termite R. flavipes [33], the fruit fly Drosophila melanogaster [34, 35], the honey bee Apis mellifera [36], the bumblebee Bombus ignites [37], the red harvester ant Pogonomyrmex barbatus [38], the ant Pheidole pallidula [39] and the fire ant Solenopsis invicta [40]. In solitary D. melanogaster, foraging genes can influence a number of behavioral traits, which can also be modified by the social context [41-43]. However, how the complex interactions of inherent genes and social contexts give rise to variations in foraging behavior in termites is still unclear.

The subterranean termite Odontotermes formosanus (Shiraki), which is a fungus-cultivating higher termite, can construct large underground cavities and damage many kinds of trees [44, 45]. Workers exhibit large foraging areas and long foraging distances in the termite O. formosanus [46]. They need to build mud lines during their foraging to avoid adverse environmental factors and increase foraging resources [47-49]. Thus, sufficient energy reserves appear to be an important factor affecting the likelihood of foraging success in termites [50]. Isocitrate dehydrogenase (IDH) is an important rate-
limiting enzyme that catalyzes the decarboxylation of isocitrate to alpha-ketoglutarate in the tricarboxylic acid cycle (TCA) and generates nicotinamide-adenine dinucleotide (NADH), which feeds into oxidative phosphorylation to generate adenosine triphosphate (ATP) [51]. A recent study showed that IDH downregulation can disrupt active immunization against the entomopathogenic fungus Metarhizium anisopliae in the termite R. chinensis [52]. In this study, we wanted to determine whether IDH gene knockdown impairs energy metabolism involved in glucose decomposition and ATP production and further affects foraging behavior in termites. Here, we cloned the IDH gene encoding the α subunit of IDH and observed its expression pattern in O. formosanus. Subsequently, we silenced the IDH gene and explored the effect of energy metabolism alterations in the whole organism and brain on the foraging behavior of O. formosanus. Moreover, we carried out orthogonal experiments to analyze the changes in IDH-mediated foraging behavior in different social contexts (with or without Leptogenys kitteli predator ants or/and nestmate soldiers) according to four experimental parameters (velocity, distance moved, frequency and cumulative duration in food zones) in O. formosanus.

**Results**

Cloning, expression and RNAi knockdown efficiency of the IDH gene in O. formosanus
The complete open reading frame (ORF) of IDH was 1,074 bp and encoded a predicted protein of 357 amino acids (aa), which is highly conserved in social organisms (Fig 1A). The prediction of the secondary structure of the IDH protein demonstrated that the amino acid sequence contained 47.90% alpha-helices, 10.36% extended strands, 6.16% beta-turns, and 35.57% random coils (Fig 1B). The analysis of the conserved domains with the NCBI tool indicated that IDH presented the specific hit Iso_dh (26-350 aa) (Fig 1B). The evolutionary tree (Fig 1C) showed that IDH from *O. formosanus* was clustered with that from the German cockroach, *Blattella germanica*, and two termite species, *Zootermopsis nevadensis* and *Cryptotermes secundus*.

Our results showed that IDH expression in the abdomen and head was significantly higher than that in the thorax (Fig 1D, Tukey’s HSD test, $F = 30.380$, $df = 2,15$, $p < 0.001$, $n = 6$). The expression level of IDH was significantly decreased by 47.44% 3 d after the introduction of RNAi (Fig 1E, paired $t$-test, $t = -11.055$, $df = 8$, $p < 0.001$, $n = 9$). Semiquantitative RT-PCR analysis also showed that the expression of IDH was successfully inhibited (Fig 1E). In addition, there were no significant differences in survival between dsIDH-injected individuals and dsGFP-injected individuals (Fig 1F, paired $t$-test, $t = -0.419$, $df = 5$, $p = 0.692$, $n = 6$).

**IDH knockdown reduced the energy supply but increased glucose accumulation**

The IDH enzyme is an important rate-limiting enzyme in the TCA and can catalyze the decarboxylation of isocitrate to alpha-ketoglutarate [51]. Therefore, we silenced *IDH*
to investigate the changes in the NAD$^+$-IDH reaction in the TCA and the level of glucose 3 d after the injection of dsIDH. The results showed that IDH activity (Fig 2A, Wilcoxon test, $Z = -2.023$, $p = 0.043$, $n = 6$), ATP levels (Fig 2B, Wilcoxon test, $Z = -2.666$, $p = 0.008$, $n = 9$) and NADH levels (Fig 2C, Wilcoxon test, $Z = -2.201$, $p = 0.028$, $n = 6$) were significantly decreased in dsIDH-injected termites compared to dsGFP-injected termites. However, the glucose level was significantly increased compared with that in dsGFP-injected termites (Fig 2D, Wilcoxon test, $Z = -2.192$, $p = 0.028$, $n = 9$).

Silencing IDH decreased walking activity but increased foraging success

To investigate the relationship between energy metabolism and foraging behavior, we silenced IDH by injecting dsIDH and then observed the phenotypic changes in the foraging behavior (Fig 3A) of O. formosanus. We found that the foraging trajectories of dsGFP-injected individuals were denser than those of dsIDH-injected individuals (Fig 3B). Both the distance moved (Fig 3C, paired $t$-test, $t = -2.502$, df = 9, $p = 0.034$, $n = 10$) and velocity (Fig 3D, paired $t$-test, $t = -2.448$, df = 9, $p = 0.037$, $n = 10$) were significantly decreased in dsIDH-injected individuals compared to dsGFP-injected individuals. However, the frequency (Fig 3E, paired $t$-test, $t = 2.890$, df = 9, $p = 0.018$, $n = 10$) and the cumulative duration (Fig 3F, paired $t$-test, $t = 2.344$, df = 9, $p = 0.044$, $n = 10$) in food zones were significantly increased in dsIDH-injected individuals compared with dsGFP-injected individuals.
The effect of IDH on brain energy metabolism in different social contexts

We successfully obtained the brain tissue of worker termites (Fig 4A) and found that there were a large number of tracheae in the brain of *O. formosanus*. Immunocytochemistry analysis of synapsin revealed the major medullary structures in the workers’ brains (Fig 4B). Our results showed that IDH expression in the worker brain was significantly decreased 3 d after dsIDH injection (Fig 4C, Wilcoxon test, \( Z = -2.201, p = 0.028, n = 6 \)). In three different social contexts, IDH activity in the brains of dsIDH-injected worker individuals was significantly higher than that in dsGFP-injected worker individuals, respectively (Fig 4D, Wilcoxon test, \( Z_1 = -1.992, p_1 = 0.046, n_1 = 6; Z_2 = -2.201, p_2 = 0.028, n_2 = 6; Z_3 = -2.201, p_3 = 0.028, n_3 = 6 \)), while ATP levels in the brains of dsIDH-injected worker individuals were significantly lower than that in dsGFP-injected worker individuals, respectively (Fig 4E, Wilcoxon test, \( Z_1 = -2.201, p_1 = 0.028, n_1 = 6; Z_2 = -2.201, p_2 = 0.028, n_2 = 6; Z_3 = -2.201, p_3 = 0.028, n_3 = 6 \)). Additionally, we found that IDH activity in the worker brain was significantly higher in the social context with ants and soldiers than that in the social context without ants and soldiers (Fig 4D, Wilcoxon test, dsIDH: \( Z = -1.992, p = 0.046, n = 6; \) dsGFP: \( Z = -2.201, p = 0.028, n = 6 \)). However, ATP levels in the worker brain were significantly lower in the social context with ants and soldiers than that in the social context without ants and soldiers (Fig 4E, Wilcoxon test, dsIDH: \( Z = -2.201, p = 0.028, n = 6; \) dsGFP: \( Z = -2.201, p = 0.028, n = 6 \)).
The influence of IDH on foraging behavior in different social contexts

To further explore the impact of IDH on the foraging behavior of O. formosanus in different social contexts, we set up nine treatment groups with or without predator ants or/and nestmate soldiers using orthogonal experimental design L₀(3³) (Table 1, Fig 5A-D). Our results showed that the foraging trajectories of O. formosanus in the nine treatment groups varied, which was associated with the status of worker energy metabolism and different social contexts (Fig 5E). The foraging trajectories of water-injected workers became increasingly complex and chaotic along with the increase in the number of ants (from Treatment 1 to 3). The foraging trajectories of dsGFP-injected workers were similar to those of water-injected workers (from Treatment 4 to 6). However, the foraging trajectories of dsIDH-injected workers exhibited less complexity and lower flexibility when the number of ants increased (from Treatment 7 to 9).

There were significant differences in the phenotypic features of foraging behavior among the nine treatment groups in worker termites (Table 1). The velocity in Treatment 7 was much lower than that in Treatments 1, 2, 3 and 6 (Tukey’s HSD test, F = 3.433, df = 8, 99, p = 0.002). At the same time, the distance moved in Treatment 7 was extremely significantly shorter than that in Treatment 1 and significantly shorter than that in Treatments 2, 3 and 6 (Tukey’s HSD test, F = 3.964, df = 8, 99, p < 0.001).

In addition, the distance moved in Treatment 8 was significantly shorter than that in Treatment 1 (Tukey’s HSD test, F = 3.964, df = 8, 99, p < 0.001). However, the frequency in food zones in Treatment 5 was significantly less than that in Treatment 1,
and the frequency in food zones in Treatment 8 was extremely significantly lower than that in Treatment 1 and was markedly less than that in Treatment 2 (Tukey’s HSD test, $F = 4.027$, df = 8, 99, $p < 0.001$). Additionally, the cumulative duration of visits in food zones in Treatment 8 was significantly shorter than that in Treatment 1, and the cumulative duration of visits in food zones in Treatment 9 was also extremely significantly shorter than that in Treatment 1 (Tukey’s HSD test, $F = 3.095$, df = 8, 99, $p = 0.004$).

The impact degrees of three factors on the velocity and distance moved were as follows: $IDH$ expression $> \text{soldier number} > \text{ant number}$, but the impact degrees of the three factors on the frequency and cumulative duration in food zones were as follows: gene expression $> \text{ant number} > \text{soldier number}$. Among these factors, $IDH$ expression showed a significant influence on the velocity and distance moved, and frequency and cumulative duration in food zones (GLM, all $p < 0.001$). Additionally, ant number exerted a significant influence on the frequency and cumulative duration in food zones (GLM, $p_1 = 0.024$; $p_2 = 0.029$).

**Discussion**

The abnormal energy metabolism of the whole organism mediated by the IDH gene seriously altered the foraging behavior of *O. formosanus*. The *IDH* gene, associated with energy metabolism, was highly expressed in the abdomen and head of foraging workers, which supports the notion that the movement requires much energy [53],...
particularly for foraging behavior, which requires integration across different sensory modalities in worker termites. In our study, silencing *IDH* disrupted mitochondrial metabolism, including the disruption of the NAD⁺-IDH reaction in the TCA cycle and the reduction of ATP levels, NADH levels and IDH activity (Fig 2A, B and C), which caused decreased walking activity, including shorter collective distances moved and slower velocities. Additionally, the trajectories of worker termites showed lower mobility (Fig 3B). These results suggest that mitochondrial metabolism plays a vital role in the regulation of walking activity. Normally, glucose can be decomposed and then produce ATP through the glycolysis pathway and TCA cycle [51]. However, a significant increase in glucose content was found in this study after silencing *IDH*, which was also recently described in association with the downregulation of human *IDH3α* [54] and knockdown of termite *IDH* [52]. Our results indicated that silencing *IDH* disrupted the process of glucose metabolism and resulted in the accumulation of glucose [52]. Surprisingly, knocking down *IDH* increased the frequency and cumulative duration in food zones within a certain amount of foraging time, suggesting that termite colonies could ensure the food supply by increasing foraging success [35] and prolonging the time of food acquisition to reduce the negative effects of energy deficiency. The changes in the energy metabolism of the whole organism caused by *IDH* knockdown in this study provided direct evidence of the importance of the rate-limiting enzyme IDH gene in regulating metabolic homeostasis and energy supply, which is closely related to foraging behavior in termites.
IDH expression levels might change the energy metabolism pathway of the termite brain, which is energetically demanding as well as a key regulator of energy metabolism [55]. In this study, IDH knockdown significantly reduced ATP levels in the brains of worker termites but dramatically improved IDH activity. These results suggested that the production efficiency of ATP decreased. Usually, glucose is metabolized via glycolysis and the TCA cycle to generate sufficient ATP through the process of oxidative phosphorylation [56]. However, IDH downregulation disrupts the TCA cycle and thereby reduces the generation of ATP. On the other hand, it has been demonstrated that a decrease in the expression of IDH3α converts the metabolic mode from oxidative phosphorylation to aerobic glycolysis in cancer-associated fibroblasts [54]. However, glycolysis is far less efficient than the TCA cycle coupled to oxidative phosphorylation in producing ATP [56]. Therefore, the decreasing ATP levels in the brain of dsIDH-injected termites suggests that the brain might choose an inefficient way of producing ATP, such as aerobic glycolysis, resulting in a low efficiency of ATP generation. Actually, ATP is necessary to maintain homeostasis and cell survival, and the loss of intracellular ATP may result in cell necrosis or apoptosis [56, 57]. In addition, energetic constraints play a major role in neural plasticity and brain health [58]. Thus, an increase in IDH activity might reduce the damage caused by ATP deficiency in dsIDH-injected termites.

Energy metabolism in the termite brain is not only impacted by IDH expression levels but is also influenced by the social context. Some researchers have suggested that
social information can change gene expression in the brain to influence behaviors in social insects [59]. We also found that social information could affect foraging behavior by changing brain energy metabolism in termites. When worker termites are exposed to predation risks during foraging, they may exhibit an aggressive state and consume much energy [29]. At this moment, ATP levels decreased remarkably, but the IDH activity increased significantly in the brains of dsIDH- and dsGFP-injected foragers in the social context with predators, suggesting that the degree of aerobic glycolysis might be enhanced, as reported for the metabolic pathway in the brain of aggressive honey bees [57]. In other words, the brains of worker termites may prefer an inefficient but faster pathway for ATP production [56] to meet the high energy demands of foraging behavior when they are threatened by predators. On the other hand, the addition of nestmate soldiers during foraging may enhance the group cognition involved in the emergent aggregation behavior of workers and social networks [60, 61], which is energetically costly in brains [12]. Therefore, the decrease in ATP became more significant, and the increase in IDH activity was greater when soldiers and ants were present concurrently. In addition, the social context may regulate the expression of genes associated with encoding neuropeptides in the brain, such as gonadotropin-releasing hormone (GnRH), vasotocin (VT) and vasopressin (VP) [62, 63]. The sensory and higher-order integrative processing mechanisms that are used when insects behave in complex social contexts are also associated with nervous systems of the brain [64], and the energetic basis of these behaviors is a bridge between behavioral ecology and
neuroscience [16]. Moreover, intermediates of energy metabolism can impact the concentration of neurotransmitters by regulating the synthesis of precursors such as alpha-ketoglutarate, which is the precursor of the excitatory neurotransmitter glutamate produced via the TCA cycle [57, 65]. Thus, the changes in energy metabolism in the brain caused by IDH downregulation and social context might alter the energetic and nervous system bases of foraging behavior in termites.

The abnormal energy metabolism in the whole organism and the brain caused by IDH downregulation and the social context could modulate foraging behavior in termites. Silencing IDH reduced the walking activity of worker termites and simplified their foraging trajectory (Treatments 7, 8 and 9 in Table 1). This reduction in walking activity decreases significantly when the number of nestmate soldiers was increased but no ants were present (Treatment 7 in Table 1) compared to the water- or dsGFP-injected workers, with normal energy metabolism (Treatments 1, 2, 3 and 6 in Table 1), indicating that worker termites may choose a more profitable foraging strategy, such as reducing their foraging area, to relieve the energy shortage [66, 67] because more energy may be needed to cover a larger movement space [68, 69]. The foraging success of dsIDH-injected workers significantly increased (Fig 3E and F), while it decreased sharply when predator ants were present without nestmate soldiers (Treatments 8 in Table 1), indicating that predator ants seriously affect foraging behavior and that the absence of nestmate soldiers means that foraging workers face a greater risk of predation if worker termites forage for a long time and feed on a large amount of food.
when the energy supply is lacking [2]. Although energy deficiency caused the foraging workers to significantly reduce the cumulative duration of visits in food zones, the increasing number of soldiers enhanced the frequency in food zones appropriately (Treatment 9 in Table 1), illustrating that soldiers can provide social buffering to cover the damage caused by the increase in ants during foraging [24, 32, 70].

However, the foraging trajectory of worker termites with normal energy metabolism became complex when ants were present (Treatments 3 and 6 in Table 1), indicating that the increase in ants led to more purposeless walking, which is energetically demanding [19, 22]. In addition, the walking activity and foraging success of the worker termites showed no significant changes as the number of soldiers was increased together with the number of ants (Treatments 1, 2 and 3 in Table 1). Therefore, the presence of nestmate soldiers is crucial to reduce the purposeless walking of foraging workers and relieve the predation pressure from ants to help foragers to accurately find food sites [24]. Even though the increase in nestmate soldiers could decrease the purposeless walking caused by a small number of ants, foraging success still decreased when the number of soldiers increased (Treatment 5 in Table 1). In addition, orthogonal experiments showed that \( IDH \) expression extremely significantly impacted both walking activity and foraging success in foraging workers (Table 1), but predator ants only significantly influenced foraging success in foraging workers (Table 1), strongly suggesting that the inherent gene plays the dominant role in modulating foraging behavior in termites compared to the external social context [35].
In summary, we confirmed that the metabolic gene IDH was an important regulator of foraging behavior in termites, which can be influenced by different social contexts. IDH downregulation reduced ATP levels and IDH activity, and disrupted the NAD$^+$-IDH reaction in the TCA in the whole organism but increased IDH activity and decreased ATP levels in the brains of termites (Fig 6). These alterations of the energetic metabolism resulted in decreased walking activity and increased foraging success, suggesting the important role of energy metabolism in the regulation of foraging behavior (Fig 6). In addition, social contexts can also change termite foraging behavior by adding predation risk from predator ants and providing social buffering from nestmate soldiers. The foraging success of dsIDH-injected workers decreased significantly with the addition of predator ants. The number of nestmate soldiers could improve the foraging success of dsIDH-injected workers by providing social buffering to relieve the negative effect of predator ants. Regardless of how the external social context changed, the impact level of the inherent IDH gene was always highest on termite foraging behavior (Table 1), which further demonstrates that the inherent gene plays a dominant role in regulating foraging behavior in termites. Hence, the present study strongly supports the hypothesis that abnormal energy metabolism can alter foraging behavior in termites in different social contexts. Certainly, whether the change in brain energy affects neural functions that may be related to foraging behavior needs to be resolved in the future. This study enriches the understanding of the energetic basis and strategy of foraging behavior in termites by verifying an energetic alteration.
impacting foraging behavior from the whole-organism level to the brain level in different social contexts, which will be beneficial for clarifying the role of the energy supply, allocation and consumption in determining the diversity, plasticity and evolution of social behaviors in insect societies.

**Materials and Methods**

**Experimental termites**

We collected termite samples from 26 *O. formosanus* colonies in the field (Table S1). Samples of ants were collected as predators from a single *L. kitteli* colony in the field. All the termite and ant colonies were collected from Shizi Hill, Wuhan City, China. They were raised under controlled laboratory conditions (darkness, 25 ± 1 °C, 80 ± 5% relative humidity). Healthy termite and ant samples were chosen for the subsequent experiments.

**Cloning and sequencing of the IDH gene**

The termite samples were frozen in liquid nitrogen immediately after collection. RNA was extracted with RNAiso Plus (Takara Bio Inc., Tokyo, Japan), and cDNA was synthesized using a PrimeScript RT reagent kit with gDNA Eraser (Takara Bio Inc.). Gene-specific primers for the complete ORF of *IDH* were designed based on partial sequences of Unigene 32825 obtained from the transcriptome data of the heads of *O. formosanus* workers [48]. The nested polymerase chain reaction (PCR) amplification
reactions were carried out as follows: 95 °C for 4 min; 38 cycles of 95 °C for 30 s, 55 °C for 30 s and 72 °C for 80 s; and one cycle at 72 °C for 5 min. Following PCR amplification, the IDH fragment was cloned, purified and sequenced.

**Sequence and phylogenetic analyses of the IDH gene**

Sequence alignment and analysis were performed using the BLAST service of the National Center for Biotechnology Information (NCBI, [http://www.ncbi.nlm.nih.gov](http://www.ncbi.nlm.nih.gov)). Protein multiple alignment analyses were performed using MEGA 6.0 and GeneDoc 2.0 software. The deduced amino acid sequence of IDH was aligned with its corresponding orthologs from the harvester ant *Pogonomyrmex barbatus* (XP_011645897.1), the honey bee *Apis mellifera* (XP_006564183.2) and *Homo sapiens* (NP_005521.1). cDNA and amino acid sequence similarity searches were performed using the BLAST algorithm ([https://blast.ncbi.nlm.nih.gov/Blast.cgi](https://blast.ncbi.nlm.nih.gov/Blast.cgi)). The phylogenetic relationships of IDH and the homologous genes in other species were analyzed using the neighbor-joining (NJ) method with MEGA 6.0 software with 1,000 bootstrapping replicates. The prediction of protein secondary structure was performed using the Self-Optimized Prediction Method with Alignment algorithm of the online software PRABI-Lyon-Gerland ([https://prabi.ibcp.fr/htm/site/web/home](https://prabi.ibcp.fr/htm/site/web/home)).

**IDH transcription in different tissues of *O. formosanus***

The head, abdomen and thorax tissues were dissected from *O. formosanus* workers under low-temperature conditions. Thirty individual thorax specimens and 15
individual head and abdomen specimens were prepared for each replicate. RNA extraction and cDNA synthesis from these body regions were carried out as described in the section on the cloning and sequencing of the IDH gene. Quantitative real-time PCR (qRT-PCR) was performed with Hieff™ qPCR SYBR® Green Master Mix (Yeasen, China) in a QuantStudio 6&7 Flex Real Time PCR System (Applied Biosystems, Life Technologies, Milan, Italy). The relative expression levels of IDH among the three body regions were calculated using the 2^{-ΔΔCt} method [71]. Six biological replicates were performed for the RT-qPCR analysis of IDH. The primers used for RT-qPCR are listed in Table S2.

RNA interference with IDH transcription

The template cDNAs were amplified by using PCR primers that had T7 RNA polymerase sequences appended to their 5’ ends. The PCR primers for the dsRNA template (shown 5’ to 3’) are shown in Table S2. Injection was performed using a sterilized microinjector (SYS-PV820, World Precision Instruments, USA). Approximately 3 μg of dsRNA targeting IDH or GFP, or 150 nl of water was injected into the side of the thorax [72]. After injection, the termites were transferred to petri dishes with filter paper containing a 7.5% glucose solution. Individuals and brain tissues were collected separately at 3 d after injection for subsequent experiments. qPCR was performed for each sample using a QuantStudio 6&7 Flex Real Time PCR System. Then, cDNA to be used as the template for qRT-PCR was synthesized from
the total RNA of each of six individuals at 3 d after injection. Gene-specific primers were designed by using NCBI primer-BLAST and are presented in Table S2. mRNA levels were quantified using β-actin and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) as the reference genes. The relative expression levels of specific genes were calculated via the 2^(ΔΔCt) method [71]. Each treatment consisted of at least 6 biological replicates.

**Brain tissue anatomy and immunocytochemistry with synapsin**

Termite brains were dissected out in an ice-cold sterilized 0.9% sodium chloride solution in a dissecting dish cooled on ice [73] and then immediately frozen with liquid nitrogen for IDH expression and metabolite assays. The insects were anesthetized by cooling on ice. The brains were dissected out and fixed in a 4% paraformaldehyde solution in phosphate-buffered saline (PBS; in mM, 684 NaCl, 13 KCl, 50.7 Na_2HPO_4, 5 KH_2PO_4, pH 7.4) for 2 h at room temperature. After fixation, the brains were rinsed in PBS for 4 × 15 min. To minimize nonspecific staining, the brains were preincubated in 5% normal goat serum (Sigma, St. Louis, MO) in PBS containing 0.5% Triton X-100 (PBSX; 0.1 M, pH 7.4) for 3 h at room temperature. Then, the brains were incubated in the primary antibody SYNORF1 at 1:100 in PBSX at 4 °C for 3 d. After incubation, the brains were rinsed in PBS for 4 × 15 min before being incubated in the Cy2-conjugated anti-mouse secondary antibody (dilution 1:300 in PBSX; Invitrogen, Eugene, OR), at 4 °C for 1 d. The brains were finally rinsed for 4 × 15 min in PBS,
dehydrated in an ascending ethanol series, and mounted in methyl salicylate [74]. A confocal laser scanning microscope (LSM 510, META Zeiss, Jena, Germany) was used to scan the brain and to obtain serial optical images with excitation by a 488 nm laser.

**Metabolite assays**

Three days post-injection, termite individuals were weighed and then immediately crushed in liquid nitrogen and dissolved at a ratio of 10 mg of body fresh weight to 100 μL of the solution recommended by the manufacturer. Termite brains were dissected 15 min after foraging observations in different social contexts and then dissolved in the solution recommended by the manufacturer. The determination of the ATP (n = 9 replicates), NADH (n = 6 replicates), IDH (n = 6 replicates) and glucose (n = 9 replicates) contents of the whole organism and the ATP (n = 6 replicates) and IDH (n = 6 replicates) contents of the brain were performed according to the protocols provided by the manufacturers (Beyotime Biotechnology (Shanghai, China), Nanjing Jiancheng Bioengineering Institute (Jiangsu, China) and Zeye Biological Technology (Shanghai, China)). The protein concentration of the brain was determined according to the protocol of the BCA (bicinchoninic acid) protein concentration determination kit from Beijing Dingguo Changsheng Biotechnology (Beijing, China).

**Termite behavioral apparatus**

The circular behavioral apparatus included one inner ring (D = 300 mm) in which 10 termites tested and a 10 mm outer ring containing the predator ant *L. kitteli*. Both rings
of the testing arena had a 10 mm-deep wall edge (Fig 3A). The wall of the inner ring included 40 evenly spaced 1 mm slits that allowed the transmission of chemical cues and antennal contacts but were too narrow for lethal interactions [24]. Four pieces of 20 mm circular filter paper moistened with 7.5% glucose solution were placed on marked food patches.

Foraging assays

In the foraging assays performed without regard to the social context (L. kitteli predator ants and nestmate soldiers), 10 workers injected with dsIDH or dsGFP were placed in different 35 mm Petri dishes before the start of the experiment. The foraging assay began after the ten workers were added into the center of the inner ring of the testing arena with moistened filter paper, and the foraging behavior of the 10 workers was then recorded by using the trajectory tracking software EthoVision XT (Noldus Information Technology). After recording for 15 min, the tested workers were removed and then quickly placed in liquid nitrogen. Considering the adaptation period of termites in the behavioral apparatus, the features of the foraging behavior of the ten workers were analyzed after 5 min of recording.

In the foraging assays performed in different social contexts (with or without L. kitteli predator ants or/and nestmate soldiers), workers and soldiers that did not receive any injection were individually marked with red color on the abdomen by using PX-21 uni-Paint markers prior to the assay so that each individual could be identified. To reduce
the potential for injury, each body part of an individual was only marked once [24]. For
the convenience of experimental operation, different injected termites were placed in
different 35 mm Petri dishes before the start of the experiment. For example, the 24
dsIDH-injected termites were transferred to three 35 mm Petri dishes, and each Petri
dish contained eight termites. Similarly, the 24 termites that were injected with dsGFP
or water were transferred to Petri dishes respectively. Each Petri dish was covered with
filter paper moistened with water. According to orthogonal experimental design L_{9}(3^{3})
(Table 1), we transferred one or two marked workers or/and marked soldiers that were
not subjected to dsRNA or water injection separately into nine Petri dishes (Fig 5A-D).
Finally, there were a total of 10 termites in each Petri dish, including unmarked and
marked individuals.

At the beginning of the experiments, the inner ring of the testing arena was covered
with moistened filter paper. According to the orthogonal experimental design L_{9}(3^{3})
(Table 1), we placed different numbers of L. kitteli predator ant workers in the outer
ring of the testing arena for 30 min before adding termites. The foraging assay began
after the ten termites (including workers and/or soldiers) were added to the center of
the inner ring of the testing arena with moistened filter paper. The features of the
behavioral phenotypes of the 10 termites were recorded by using EthoVision XT
software. After recording for 15 min, the tested termites were removed and then quickly
placed in liquid nitrogen. The features of the foraging behavior of eight workers were
analyzed after 5 min of recording, which was also the procedure for the foraging assay without the social context described above.

Four important phenotypic parameters of foraging behavior were measured, including the velocity and distance moved, and the frequency and cumulative duration in food zones. In this study, the velocity and distance moved were used to describe the walking activity of the worker termites, and the frequency and cumulative duration in food zones were used to describe the foraging success of the worker termites [35, 75].

**Statistical analysis**

The foraging behaviors of termites were monitored using a digital camera (BASLER, acA1920-40gc) coupled to video tracking software (EthoVision XT 14.0, Noldus Information Technology). All statistical analyses were conducted using IBM SPSS Statistics 19.0 software. The Shapiro-Wilk test was used to verify whether the data conformed to a normal distribution or not. Most of the data showed a normal distribution, except for those for the RNAi efficiency in the brain and the metabolite assays. The expression pattern of IDH and the phenotypic parameters of foraging behavior in the orthogonal experiments were analyzed with Tukey’s HSD test. Gene expression, RNAi efficiency in the whole-organism and foraging assays after silencing IDH without regard to the social context were analyzed with paired t-tests. The impact degrees of three factors (gene, ant number and soldier number) on worker foraging behavior were determined with a general linear model (GLM). The assessment of RNAi
efficiency in the brain and metabolite assays was performed with the Wilcoxon test.

The significance level in this study was set at $p < 0.05$.

Acknowledgments

We would like to thank Dr. Zhao Xincheng from Henan Agriculture University for technical help with the brain tissue anatomy and immunocytochemistry of termites.

Funding

Funding for this research was supported by the Fundamental Research Funds for the National Natural Science Foundation of China (grant number: 31772516).
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Fig 1. Cloning, expression and RNAi efficiency of the IDH gene in *O. formosanus*. (A) Multiple alignments of the amino acid sequences deduced for IDH in *O. formosanus* and three other social species. Identical residues were shaded in black. Dashed lines indicated the gaps. (B) Secondary structure prediction of the amino acid sequence of IDH and conserved domains of IDH. Random coils were shown in purple, alpha-helixes were shown in blue, beta-turns were shown in green, and extended strands were shown in red. The gene showed an Iso_dh specific hit (26-350). (C) Phylogenetic relationships of the IDH proteins from *O. formosanus* and 11 other social species. (D) Expression patterns of IDH in the head, thorax and abdomen of workers. (E) Expression level and semiquantitative detection of IDH 1, 3 and 4 days after dsRNA injection in workers; (F) Survival 3 d after dsRNA injection in workers. Different lowercase letters in a column indicate significant differences by Tukey’s HSD test. ***, p < 0.001, n.s. means no significant difference.
Fig 2. Alterations in energy metabolism after silencing IDH in O. formosanus. (A) IDH activity in workers 3 d after dsIDH injection; (B) ATP levels in workers 3 d after dsIDH injection; (C) NADH levels in workers 3 d after dsIDH injection; (D) Glucose levels in workers 3 d after dsIDH injection. * p < 0.05.
Fig 3. Silencing IDH decreased walking activity but increased foraging success in O. formosanus.

(A) Behavioral apparatus (1, camera; 2, testing arena). (B) Foraging trajectories of dsIDH-injected (deep green color) and dsGFP-injected (deep blue color) workers. (C) The distance moved of dsIDH-injected and dsGFP-injected workers. (D) The velocity of dsIDH-injected and dsGFP-injected workers. (E) The frequency in food zones of dsIDH-injected and dsGFP-injected workers. (F) The cumulative duration in food zones of dsIDH-injected and dsGFP-injected workers. * p < 0.05.
Fig 4. Effect of IDH silencing on brain energy metabolism in O. formosanus in different social contexts. (A) Full brain tissue of workers; (B) Confocal image of the worker brain; (C) Expression level of IDH in the brains of workers 3 d after dsIDH injection; (D) IDH activity in the brains of dsIDH-injected and dsGFP-injected workers in different social contexts. (E) ATP levels in the brains of dsIDH-injected and dsGFP-injected workers in different social contexts. * p < 0.05.
Workers without (A) or with (B) soldiers were confined in the inner ring (in gray) without ants in the outer ring (areas in light green). The wall of the center dish was cut vertically to make 1 mm-wide slits. Predation risk was perceived by the antennation of workers of the ant *L. kitteli* through the slits. The predator ants were placed in the outer ring (areas in deep orange) without (C) or with (D) soldiers. (E) Representative maps of foraging trajectories in the water-injected (in blue), dsGFP-injected (in deep blue) and dsIDH-injected (in deep green) workers. Numbers in the upper left corner of each trajectory map indicate different treatment numbers in the orthogonal experiments L₉ (3³).
Fig 6. The role of the IDH gene and the social context in regulating the foraging behavior of *O. formosanus*. IDH silencing impaired the NAD⁺-IDH reaction in the TCA, leading to a decrease in ATP levels, IDH activity and the NADH levels but an increase in glucose levels in the whole organism, resulting in increased IDH activity but decreased the ATP level in the brain. When dsIDH-injected workers foraged together, their velocity and distance moved decreased, but their frequency and cumulative duration in food zones increased, suggesting that IDH downregulation reduced walking activity but enhanced foraging success. The social context could also alter the brain energy metabolism of foraging workers, including decreasing ATP levels but the increasing IDH activity in the social context with ants and soldiers, which further changed the foraging behavior of the workers. When predator ants were present, the dsIDH-injected workers decreased their frequency and cumulative duration in food zones, showing a significant decline in foraging success. However, the increase in the number of
nestmate soldiers strengthened social buffering to relieve the negative effect of predator ants on worker foraging behavior and, thus, improved the foraging success of dsIDH-injected workers. Our orthogonal experiments verified that the role of the IDH gene as an inherent factor was dominant in modulating termite foraging behavior compared to the external social context (predator ants and nestmate soldiers). Thus, abnormal energy metabolism mediated by IDH altered termite foraging behavior in different social contexts.
Table 1 The features of foraging behavior in *O. formosanus* under different RNAi treatments and social contexts

| Treatment number | Factors (levels) | Results |  |  |  |  |  |
|------------------|------------------|---------|---------|-------|-------|-------|-------|
|                  | Gene             | Ant number | Soldier number | Velocity (cm/s) | Distance moved (cm) | Frequency in food zones | Cumulative duration in food zones (s) |
| 1                | water            | 0        | 0        | 1.62±0.10a | 900.31±56.41a       | 46.60±2.96a | 41.86±1.85a |
| 2                | water            | 10       | 1        | 1.60±0.12a | 879.71±74.34ab      | 41.53±4.02ab | 38.92±2.18ab |
| 3                | water            | 20       | 2        | 1.59±0.11a | 856.93±61.35ab      | 39.52±3.64abc | 36.61±2.27ab |
| 4                | dsGFP            | 0        | 1        | 1.45±0.13ab | 799.60±63.44abc    | 39.63±4.86abc | 38.16±2.21ab |
| 5                | dsGFP            | 10       | 2        | 1.33±0.09ab | 707.54±53.08abc    | 27.35±4.14bc | 29.72±4.11ab |
| 6                | dsGFP            | 20       | 0        | 1.57±0.12a | 859.54±70.34abc    | 38.70±4.47abc | 36.03±3.22ab |
| 7                | dsIDH            | 0        | 2        | 1.09±0.09b | 583.60±48.83c      | 30.60±3.97abc | 34.30±3.10ab |
| 8                | dsIDH            | 10       | 0        | 1.22±0.07ab | 630.35±48.70bc     | 22.67±2.75c  | 27.72±2.46b  |
| 9                | dsIDH            | 20       | 1        | 1.31±0.10ab | 663.02±59.63abc    | 31.22±3.29abc | 29.13±2.88b  |

Velocity (cm/s)

- **R** indicates the impact degree of the three factors on the features of foraging behavior obtained from the L$_9$ (3$^3$) orthogonal experiment. A higher **R** value for a factor suggests a stronger effect on foraging behavior. The data in the table are the mean ± SEM, and different letters in a column indicate significant differences by Tukey’s HSD test (**p** < 0.05, **n** = 12). Asterisks indicate significant differences by GLM (**** < 0.001, 0.01 < ** < 0.05).
## Supporting information

### Table S1 Distribution of the 26 O. formosanus colonies for each experiment*

| Experiments                                      | Replicates | Colonies          |
|--------------------------------------------------|------------|-------------------|
| Expression pattern of IDH                        | 6          | 1,2,3             |
| RNAi efficiency for IDH                          | 9          | 4,5,6             |
| Survival analysis for IDH                        | 6          | 5,6               |
| Evaluation of IDH enzyme activity                | 6          | 7,8               |
| Evaluation of ATP level                          | 9          | 7,8,9             |
| Evaluation of NADH level                         | 6          | 10,11             |
| Evaluation of glucose level                      | 9          | 10,11,12          |
| Evaluation of IDH enzyme activity in the brain   | 6          | 13,14,15          |
| Evaluation of ATP level in the brain             | 6          | 13,14,15          |
| RNAi efficiency for IDH in the brain             | 6          | 16,17             |
| Foraging behavior for IDH                        | 10         | 17,18,19,20       |
| Orthogonal experiments for foraging assays       | 12         | 21,22,23,24,25,26 |

* All the termite samples were collected from 26 colonies of *O. formosanus* on Shizi Hill, Wuhan City, Hubei Province, China.
| Gene Name                          | Abbreviation | Orientation | Primer Sequences                          |
|-----------------------------------|--------------|-------------|-------------------------------------------|
| Isocitrate Dehydrogenase dsRNA    | dsIDH        | Forward     | GGATCCTAATACGACTCACTATAGGGACCCTGTCGATC    |
|                                   |              | Reverse     | ACTGGAAGGGGACCTGACCTGAGAAGCGGAAG          |
| Green Fluorescent Protein dsRNA   | dsGFP        | Forward     | GGATCCTAATACGACTCACTATAGGGCTTTATCTGCTGCT |
|                                   |              | Reverse     | GGATCCTAATACGACTCACTATAGGGTTGGCTTTATCTGCT |
| Isocitrate Dehydrogenase          | IDH          | Forward     | GAGTGGTGTCAAATGGTGCC                      |
|                                   |              | Reverse     | AAACCTGCGGCTGACCTC                       |
| beta-actin                        | β-actin      | Forward     | CTGGAGAAGTCATACGAGTTG                     |
|                                   |              | Reverse     | AGAAGGAAAGGCTGGAACA                      |
| Glyceraldehyde Phosphate Dehydrogenase | GAPDH    | Forward     | TCACTGCAACGCAGAAGACT                      |
|                                   |              | Reverse     | ATGGAAACTGGCACCACGGA                     |