
Rigid Body Dynamics Simulation Based on Graph Neural Networks with Constraints

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Abstract

In recent years, the utilization of Graph Neural Network (GNN)-based methods for simulating complex physical systems has significantly advanced the field of numerical science and engineering. Despite their success, current GNN-based methods for rigid body dynamic simulation are constrained to relatively simple scenarios, hindering their practical use in industrial settings where complex mechanical structures and interconnected components prevail. These methods face challenges in handling intricate force relationships within rigid bodies, primarily due to the difficulty in obtaining force-related data for objects in industrial environments. To address this, we propose a novel constraint-guided method that incorporates force computations into GNN-based simulations. The model incorporates computations related to both contact and non-contact forces into the prediction process. Additionally, it imposes physical constraints on the prediction process based on Kane's equations. We have rigorously demonstrated the model's rationality and effectiveness with thorough theoretical demonstration and empirical analysis.

1. Introduction

Simulation of rigid body dynamics plays a pivotal role in numerous domains within numerical science and engineering (Landau et al., 2008; Thijssen, 2007; Szabó & Babuška, 2021), including mechanical engineering (Brach, 2007), automotive design (Tong, 2000), biomechanics (Silva et al., 1997), and virtual reality (Sauer & Schömer, 1998), enabling the accurate modeling and analysis of object motion, stress, and interactions for optimization and performance enhancement. Traditional methods for simulating rigid body

systems often rely on intricate physical equations tailored to specific domains (Huston & Passerello, 1979; Mirtich, 1996; Featherstone, 2014). This necessitates a precise understanding of the physical properties of the analyzed objects and the physical environment in which they exist (Thijssen, 2007). Furthermore, such methods require a thorough grasp of relevant physical principles and relationships. With the rise of deep learning, connectionist machine learning approaches have demonstrated outstanding performance across various fields, bypassing the need for acquiring intricate prior knowledge and rules about the internal complexities of systems through a data-driven approach (Dong et al., 2021). Deep learning has simplified many complex problems in different domains and made previously unsolvable problems tractable. In recent years, the emergence of the Graph Neural Network (GNN) based physics simulation methods has opened new avenues for addressing rigid body dynamics analysis problems using deep learning (Sanchez-Gonzalez et al., 2020; Pfaff et al., 2021). Such approaches effectively extend neural network methods from traditional domains such as images, videos, and text into the realm of real-world physical analysis. Leveraging GNNs, such methods analyze the motion states of objects modeled as graphs, achieving commendable results in related methodologies.

However, the current GNN-based methods concerning rigid body dynamic analysis are still limited to analyzing relatively simple and ideal scenarios, often focusing on the analysis of common geometric shapes (Allen et al., 2023; Bhattoo et al., 2022; Sanchez-Gonzalez et al., 2020; Han et al., 2022a; Bhattoo et al., 2022; Bishnoi et al., 2023). This limitation hinders the practical application of such methods in industrial settings. In industrial environments, complex mechanical structures often involve numerous interconnected components, including intricate force relationships within rigid bodies and the resulting mutual influences between objects. Presently, GNN-based methods encounter significant challenges in addressing such problems, primarily because force-related data for objects is often challenging to observe and obtain (Iglberger & Rüde, 2011). As a result, these methods typically rely solely on positional and velocity information of objects for analysis. The difficulty in acquiring training data relevant to forces, coupled with the necessity for force relationship analysis, places these

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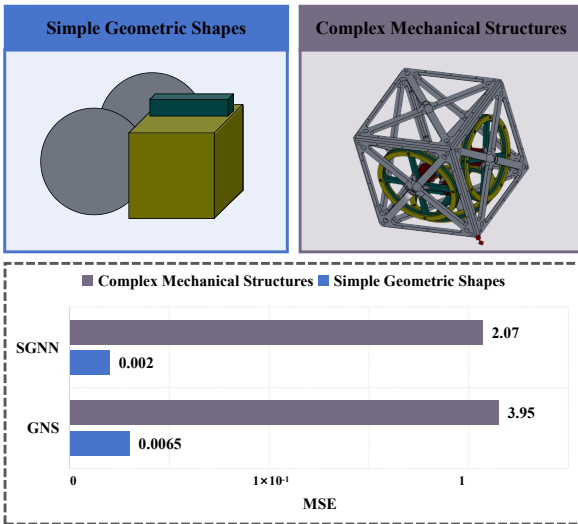


Figure 1. The comparison between the complex mechanical body scenarios we utilized and the general geometric body scenarios employed by other methods, the following illustrates the performance of two GNN-based Physical Dynamics Simulators in different scenes. From the figure, it is evident that both SGNN (Han et al., 2022a) and GNS (Sanchez-Gonzalez et al., 2020) methods exhibit significantly higher Mean Squared Error (MSE) in their predictive results for the dynamic analysis of complex mechanical structures compared to simple geometric shapes.

methods in a dilemma when applied to rigid body dynamics analysis with complex mechanical structures. The comparisons and experiments within Figure 1 validate such a conclusion, demonstrating a significant degradation in the performance of the relevant GNN-based methods for the analysis of complex mechanical structures.

To address this challenge, we endeavor to draw inspiration from traditional multibody dynamics analysis methods and incorporate relevant techniques. It is noteworthy that while GNN-based dynamic analysis methods can assist in circumventing the need for acquiring intricate prior knowledge and rules about the internal complexities of systems, the output results must adhere to the principles and rules of the physical world. Some researchers argue that a purely data-driven learning paradigm may struggle to achieve optimal results for physical problems (Bhattoo et al., 2022; Han et al., 2022a; Bishnoi et al., 2023). Therefore, we aim to integrate the principles of traditional multibody dynamics analysis with GNN-based simulators, incorporating prior constraints, to address the challenges associated with rigid body dynamics analysis in complex mechanical structures.

Building upon the aforementioned analysis, we propose a *Multibody Dynamics Guided GNN Simulator*, dubbed MDGS, to conduct a more thorough analysis of rigid body dynamics scenarios involving intricate mechanical structures. Initially, we introduce a graph modeling pattern for

efficient analysis of forces in physical systems, tailored for multibody dynamics. Subsequently, we introduce calculations related to forces, encompassing both contact and non-contact forces, into the methodology. These force calculations are performed using neural networks without the introduction of additional data beyond conventional GNN-based physical dynamics simulators. Then, the computed forces are integrated into the succeeding analyses to adapt to complex mechanical scenarios. Furthermore, we imposed constraints on the model output based on the Kane equations tailored for addressing multibody dynamics analysis problems. This ensures the accuracy and rationality of the model's computed results. The major contributions are as follows.

- We introduce a novel method, referred to as MDGS, to incorporate force computation into GNN-based simulations for the analysis of rigid body systems. This approach enhances the treatment of multibody dynamics scenarios, expanding the potential applications of such methods in industrial settings.
 - We imposed physical constraints on our method based on Kane's equations, offering a more precise analysis of rigid body dynamics. These constraints are derived from rigorous theoretical analysis and proof.
 - We provide the implementations of MDGS for multi-body dynamics simulation tasks and tested on multiple scenarios, the results consistently showcase the outstanding performance of MDGS.

2. Related Works

2.1. GNN-based Physical Dynamics Simulators

GNN-based physical dynamics simulators (Sanchez-Gonzalez et al., 2020) leverages GNNs (Scarselli et al., 2009; Kipf & Welling, 2017) to simulate predictive representations for graph-modeled physical systems. Such approaches find widespread applications in computing atomic forces (Hu et al., 2021), simulating particle-modeled and mesh-modeled physical systems (Li et al., 2019; Sanchez-Gonzalez et al., 2020; Pfaff et al., 2021; Han et al., 2022b). In recent research, some methods (Han et al., 2022a; Bhattoo et al., 2022; Bishnoi et al., 2023) have proposed incorporating prior knowledge from relevant domains into the GNN-based physical dynamics simulator to enhance its performance. This includes integrating concepts such as Subequivariant and Lagrangian equations into the model framework, making the model more closely aligned with real-world physical systems. Additionally, some approaches (Linkerh  gner et al., 2023) optimize the model by collecting data in real-world environments as supplementary information. Our method introduces force analysis between physical

entities in rigid body dynamics scenarios into the system and employs Kane's equations (Kane & Levinson, 1985) to impose constraints on the system output, ensuring a more realistic simulation performance.

2.2. Dynamics Analysis

The framework of traditional mechanical analysis relies on either force-based or energy-based approaches (Wyk et al., 2022) for rigid body dynamics analysis. Energy-based approaches involve the computation of all the unknown energy based on the equations of equilibrium and hence are cumbersome for large structures (Finzi et al., 2020). The force-based multibody dynamics model solves the forward kinematics and inverse dynamics of the main mechanisms by employing two recursive algorithms centered around the Newton–Euler formulation (Gonçalves et al., 2023). And some methods use ODEs to calculate multibody corresponding conditions to approximate an optimal solution (Schubert et al., 2023; Nada & Bayoumi, 2023; Zhang et al., 2023). We amalgamate select concepts from traditional rigid body dynamics analysis with the newly introduced GNN-based approach to achieve improved and simplified predictions.

3. Methodology

3.1. Preliminary

3.1.1. GNN-BASED SIMULATORS

GNN-based simulators predict the dynamical states of physical systems. Consider a physical system comprising M elements, collectively forming N objects. At time t , GNN-based physical dynamics simulators model the aforementioned physical system using a graph $G^{(t)} = \{\mathcal{V}^{(t)}, \mathcal{E}^{(t)}\}$, where the node set $\mathcal{V}^{(t)}$ of graph $G^{(t)}$ represents different elements constituting objects. The feature $\mathbf{z}_i^{(t)}$ of node i in the set $\mathcal{V}^{(t)}$ includes position information $\vec{x}_i^{(t)}$, velocity information $\vec{v}_i^{(t)}$, and \mathbf{k}_i representing the properties of the object to which the element belongs. In other words, $\mathbf{z}_i^{(t)}$ is defined as the concatenation of these vectors: $\mathbf{z}_i^{(t)} = [\vec{x}_i^{(t)} || \vec{v}_i^{(t)} || \mathbf{k}_i]$, where ‘||’ denotes vector concatenation along the first dimension. $\mathbf{Z}^{(t)} = \{\mathbf{z}_i^{(t)}\}_{i=1}^M$ denotes the set of all $\mathbf{z}_i^{(t)}$. The edge set $\mathcal{E}^{(t)}$ of graph $G^{(t)}$ characterizes the connectivity relationships between elements. For instance, some approaches (Han et al., 2022a; Bhattoo et al., 2022) judge that if the distance between element i and j is less than a certain threshold, the edge (i, j) is connected. Subsequently, GNN-based simulator utilizes the GNN model $g(\cdot)$ to predict the values of $\mathbf{Z}^{(t+1)}$ based on $G^{(t)}$, such process can be formalized as follows:

$$\mathbf{Z}^{(t+1)} = g(G^{(t)}). \quad (1)$$

3.1.2. KANE'S EQUATION

Kane's equation (Kane & Levinson, 1985) is a mathematical formulation used in the field of multibody dynamics to describe the motion of interconnected rigid bodies. For a nonholonomic system \mathcal{S} , the number of independently variable motions or deformations is referred to as degrees of freedom. For each degree of freedom, there exists a corresponding generalized velocity. These velocity vectors associated with generalized coordinates are typically denoted by \dot{q} , where q represents the generalized coordinates. Specifically, with respect to the generalized coordinate q_i and its corresponding generalized velocity \dot{q}_i , if \dot{q}_i is independent of the other velocities in the system, meaning that it cannot be expressed or derived from other velocities in the system's description, then \dot{q}_i is referred to as an independent velocity. Furthermore, the partial velocity $u'_{ij} = \frac{\partial \dot{q}_i}{\partial q_j}$ denotes the partial derivative of the velocity with respect to coordinate q_i when coordinate q_j is varied, within the context of a multi-coordinate system.

With the aforementioned concepts, we could define the general active force according to the γ -th independent velocity of \mathcal{S} as follows:

$$K_\gamma = \sum_{i=1}^n \vec{F}_i \cdot \vec{u}'_{i\gamma}, \quad (2)$$

where \vec{F}_i denotes the force acting on the i -th mass point in the system \mathcal{S} , and $\vec{u}'_{i\gamma}$ represents its partial velocity with respect to the γ -th independent velocity, n is the number of mass points. Likewise, we could define the generalized inertia forces according to the γ -th independent velocity as follows:

$$K_\gamma^* = \sum_{i=1}^n \left(-m_i \ddot{\vec{r}}_i \right) \cdot \vec{u}'_{i\gamma}, \quad (3)$$

where m_i represents mass, while \vec{r}_i represents the vector radius. The Kane's equation can then be formulated as follows:

$$K_\gamma^* + K_\gamma = 0, \gamma = \{1, 2, \dots, d\}, \quad (4)$$

d is the number of independent velocities.

3.2. Method

3.2.1. GRAPH CONSTRUCTION

Two commonly used object partitioning approaches in the field of physical simulation are point clouds (Gross et al., 2002) and finite elements (Pasciak, 1995), both of which have been widely applied in GNN-based simulators. Point clouds essentially involve decomposing objects into indivisible elements using point cloud construction techniques. On the other hand, the finite element method divides objects

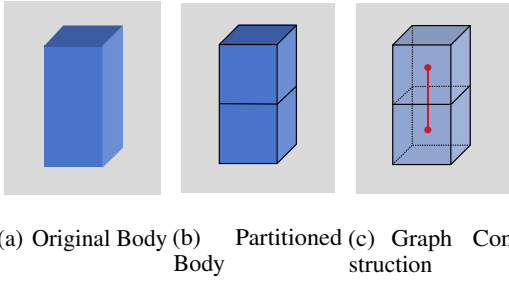


Figure 2. Illustration of the Graph Construction Process.

into individual polyhedral elements, where the edges and vertices of these elements constitute the graph for GNN-based simulators. In our approach, we integrate the above two methods by employing the finite element method to partition rigid bodies into different elements and designate the centroid of each element as a node in the graph. In the specific implementation, we assume uniform density within each element and utilize its geometric center as its centroid. We believe that this approach maximally preserves the kinematic properties of objects while aligning more closely with the commonly used analytical paradigm in multibody dynamics (Kane & Levinson, 1985). Figure 2 provides an illustrative example of our graph construction method. As illustrated in Figure 2(c), the centroids serve as nodes in the graph, and the centroids of adjacent elements are connected by edges. The specific methodology for edge construction will be elaborated in detail in Section 3.2.2.

3.2.2. CALCULATING FORCES

In the realm of multibody dynamics, the magnitude of forces acting on an object plays a crucial role in determining its state. However, due to the inherent difficulty in measuring and observing the force conditions of objects, existing GNN-based simulators do not currently incorporate force information into the prediction computation process. To address this issue, we employ neural networks to predict the forces acting on rigid bodies and integrate these predicted values into the model. Specifically, we categorize the forces acting on objects into two components: contact force and non-contact force.

For contact forces, such as frictional and collision forces, we propose the construction of a contact-oriented graph $\bar{G}^{(t)} = \{\mathcal{V}^{(t)}, \bar{\mathcal{E}}^{(t)}\}$. In $\bar{G}^{(t)}$, $\mathcal{V}^{(t)}$ represents the centroids corresponding to distinct elements, with attributes comprising the input positional and velocity information of these elements at time t . Additionally, $\mathcal{V}^{(t)}$ includes property about the particular object each element is affiliated to, providing the information concerning the whole object. $\bar{\mathcal{E}}^{(t)}$ comprises only those edges determined by the existence of contact between elements. As depicted in Figure 4(a), if the distance between the centroids of two elements, denoted as

i and j , is less than r , and they belong to different objects, we consider these elements to be in contact, and the edge (i, j) is established, r is a threshold defined by a hyperparameter. In this manner, we derive a graph $\bar{G}^{(t)}$ specifically illustrating contact relationships. Subsequently, we employ $\bar{G}^{(t)}$ to predict the force set $\bar{F}^{(t)}$, where $\bar{F}^{(t)} \in \mathbb{R}^{N \times 3}$. The i -th vector $\bar{f}_i^{(t)}$ of $\bar{F}^{(t)}$ denotes the contact forces acting upon the centroid of the i -th element. Formally, we have:

$$\bar{F}^{(t)} = g^{\bar{F}}(\bar{G}^{(t)}), \quad (5)$$

where $g^{\bar{F}}(\cdot)$ denotes a GNN specifically designed for contact force prediction. $\bar{f}_i^{(t)}$ will be set to the zero vector if i is not an endpoint of any edge in $\bar{\mathcal{E}}^{(t)}$.

For non-contact forces, such as gravity, we propose constructing the object-oriented graph $\tilde{G}^{(t)} = \{\mathcal{V}^{(t)}, \tilde{\mathcal{E}}^{(t)}\}$, where $\mathcal{V}^{(t)}$ remains identical to that of $\bar{G}^{(t)}$. As illustrated in Figure 4(b), if two elements i and j belong to the same object and share faces between them, then an edge (i, j) is established within $\tilde{\mathcal{E}}^{(t)}$. Subsequently, we predict the non-contact force set $\tilde{F}^{(t)}$. Formally, this prediction is expressed as:

$$\tilde{F}^{(t)} = g^{\tilde{F}}(\tilde{G}^{(t)}), \quad (6)$$

where $g^{\tilde{F}}(\cdot)$ denotes a GNN specifically designed for non-contact force prediction. The final force set is obtained by combining the contact and non-contact force sets:

$$F^{(t)} = \{f_i^{(t)} : f_i^{(t)} = \bar{f}_i^{(t)} + \tilde{f}_i^{(t)}\}, \quad (7)$$

where $f_i^{(t)}$ and $\tilde{f}_i^{(t)}$ represent the i -th force vector of $F^{(t)}$ and $\tilde{F}^{(t)}$ respectively.

3.2.3. MAKING PREDICTIONS

Subsequently, we predict $G^{(t+1)}$ based on both $G^{(t)}$ and $F^{(t)}$. We first attach the calculated force to the attribute z of each node. For the i -th node with attribute feature z_i , the new attribute feature z'_i is calculated as follows:

$$z'_i = [z_i || \lambda F_i^{(t)}], \quad (8)$$

where λ is a hyperparameter for controlling the magnitude of the influence of $F^{(t)}$. We denote the modified $G^{(t)}$ as $G'^{(t)}$ and calculated the prediction as follows:

$$Z^{(t+1)} = g(G'^{(t)}), \quad (9)$$

where $g(\cdot)$ denotes the GNN for state prediction. Motivated by (Allen et al., 2023), we introduced a rigid body structural constraint module in our method. This module ensures the invariance of rigid body structures in the predicted results. Specifically, we calculate the relative positions of various nodes on the object based on the initial values. Subsequently,

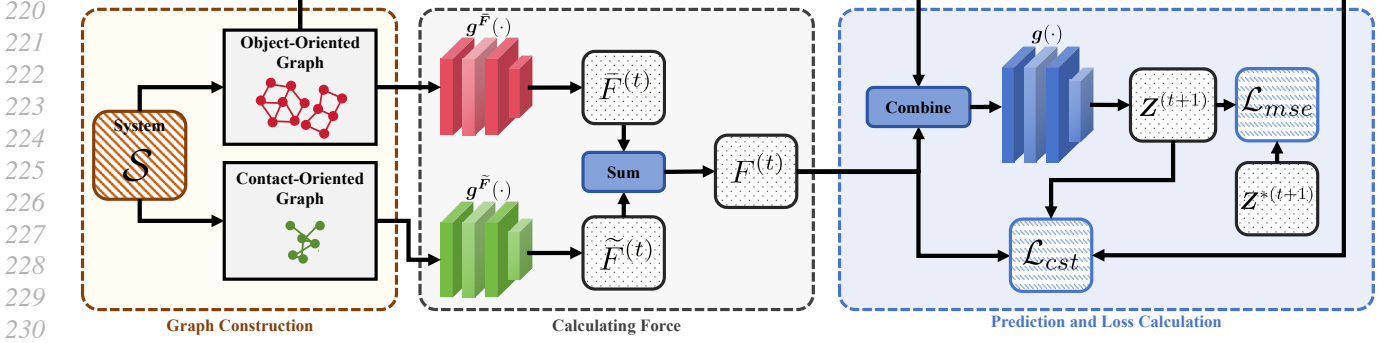
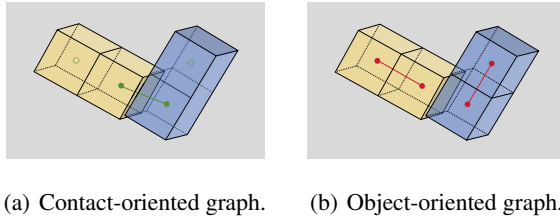


Figure 3. The framework of our proposed method. In the figure, the framework is segmented based on the stages of method execution, dividing it into three parts.



(a) Contact-oriented graph. (b) Object-oriented graph.

Figure 4. Illustration of contact-oriented graph and object-oriented graph.

we re-align our predicted results to the original relative positions, ensuring that the rigid body structure remains unchanged. A detailed description of this module can be found in [Appendix B.1](#).

$Z^{(t+1)}$ will then be utilized for calculating an MSE loss against the ground truth state $Z^{*(t+1)}$ for the backpropagation of the whole framework:

$$\mathcal{L}_{mse} = \text{MSE}(Z^{(t+1)}, Z^{*(t+1)}). \quad (10)$$

3.2.4. MODEL CONSTRAINTS BASED ON KANE'S EQUATION.

So far, we have successfully incorporated force computation to assist in predictions within the GNN-based simulators. However, such computation still faces challenges in ensuring accuracy. As discussed within section 3.1.2, Kane's equations provide a physical constraint for rigid body dynamical systems, and we aim to employ such constraints to refine our model, enhancing the analytical fidelity to physical reality.

Theoretically, a neural network capable of accurately predicting rigid body systems should adhere to the following principles:

Theorem 3.1. Consider a rigid body dynamics system denoted as S with d independent velocities. At time t , the system's state is represented as $G^{(t)}$, constructed according to the detailed graph construction procedure outlined in

[Section 3.2.1](#) and [3.2.2](#). For any Graph Neural Network (GNN) denoted as $g(\cdot)$, which demonstrates accurate prediction capabilities for $F^{(t)}$ and $Z^{(t+1)}$ based on $G^{(t)}$, the following equation must be satisfied:

$$\sum_i F_{\gamma,i}^{(t)} u_\gamma - \sum_i M_i \frac{1}{\delta} (g(G^{(t)})_{[v_\gamma,i]} - v_\gamma^{(t)}) u_\gamma + \Psi^{(t)} = 0, \forall \gamma \in \{1, 2, \dots, d\}, \quad (11)$$

where γ denote the index of independent velocity, $F_{\gamma,i}^{(t)}$ is the predicted force value of element i along γ -th independent velocity, M_i is the mass of element i , $g(G^{(t)})_{[v_\gamma,i]}$ denote the predicted $v_\gamma^{(t+1)}$ with $g(\cdot)$, δ is the time step length, $\Psi^{(t)}$ is a disturbance term directly proportional to the sum of the radii from different mass points to the center of mass in each element.

The proof can be found in [Appendix A.1](#). Theorem 3.1 furnishes us with a specific methodology for constraining the model. From theorem 3.1, it is evident that the constraints proposed in Equation 11 hold along any independent velocity. In practical implementation, we can determine the number of equations the model needs to satisfy based on the distinct independent velocities associated with different systems.

To further constrain the perturbation terms in Theorem 3.1, we propose the following theorem:

Theorem 3.2. Given the conditions within Theorem 3.1, then at time t , the following inequality holds:

$$-\tau \sum_{i=1}^n (|\vec{R}_i^{(t)}|) \omega_\gamma - \sum_{i=1}^n \frac{2}{5} M_i \tau^2 \dot{\omega}_i^{(t)} \leq \Psi^{(t)} \leq \tau \sum_{i=1}^n (|\vec{R}_i^{(t)}|) \omega_\gamma + \sum_{i=1}^n \frac{2}{5} M_i \tau^2 \dot{\omega}_i^{(t)}, \quad (12)$$

where $\vec{R}_i^{(t)}$ denotes the interval force act on element i , τ is the max radius τ among all elements, ω_γ denotes unit angular velocity value along the direction of the γ -th independent

velocity, $\dot{\omega}_i^{(t)}$ represents the max angular acceleration of element i .

Based on Theorem 3.1 and Theorem 3.2, we proposed the following loss function for the upper bound of constrain:

$$\mathcal{L}_{sup} = \text{Relu}\left(\sum_i F_{\gamma,i}^{(t)} u_\gamma - \sum_i M_i \frac{1}{\delta} (g(G^{(t)})_{[v_\gamma,i]} - v_\gamma^{(t)}) u_\gamma - \tau \sum_{i=1}^n (|\vec{R}_i^{(t)}|) \omega_\gamma - \sum_{i=1}^n \frac{2}{5} M_i \tau^2 \dot{\omega}_i^{(t)}\right), \quad (13)$$

where $F_{\gamma,i}^{(t)}$ and $|\vec{R}_i^{(t)}|$ can be calculated with force prediction, while $\dot{\omega}_i^{(t)}$ is calculated based on variant within coordinates, $\text{Relu}(\cdot)$ denote the ReLU function, M_i is calculated based on the properties of the object, the details can be found in Appendix B.1. Similarly, we can calculate the loss function for the lower bound of constrain:

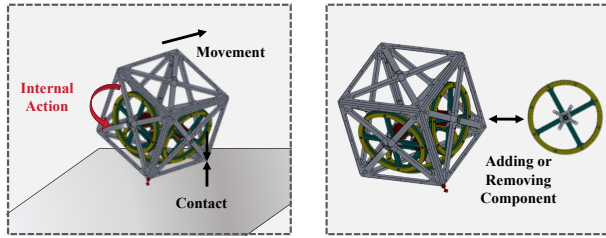
$$\mathcal{L}_{inf} = \text{Relu}\left(-\sum_i F_{\gamma,i}^{(t)} u_\gamma + \sum_i M_i \frac{1}{\delta} (g(G^{(t)})_{[v_\gamma,i]} - v_\gamma^{(t)}) u_\gamma - \tau \sum_{i=1}^n (|\vec{R}_i^{(t)}|) \omega_\gamma - \sum_{i=1}^n \frac{2}{5} M_i \tau^2 \dot{\omega}_i^{(t)}\right), \quad (14)$$

The overall loss for constrain is:

$$\mathcal{L}_{cst} = \mathcal{L}_{sup} + \mathcal{L}_{inf}. \quad (15)$$

\mathcal{L}_{cst} will only be used to update g^F and $g^{\bar{F}}$. \mathcal{L}_{cst} and \mathcal{L}_{mse} are summed up for the final loss. The overall framework is illustrated within Figure 3.

4. Experiments



(a) Our dataset incorporates interactions between complex mechanical bodies and the environment, the overall movement of mechanical bodies, and the internal actions of components within the mechanical bodies.

(b) We modify the attributes and motion patterns of mechanical bodies by adding or removing mechanical components within them. Such modifications allow us to test the performances of the algorithms across various scenarios.

Figure 5. The underlying principles behind the construction of our dataset.

4.1. Environmental Setup

4.1.1. DATASET AND SYSTEMS

To validate our methodology, we conduct comprehensive testing on diverse datasets derived from four distinct mechanical systems. Specifically, we utilized the Cubli robot (Gajamohan et al., 2012) (a metallic mechanical cube capable of performing various maneuvers using three flywheels), complex rotating bodies (intricate mechanical structures featuring components such as bearings and hinges), vehicles (mechanical cars operating under different conditions and environments), and space robots (Wang et al., 2022) (robots equipped with mechanical arms for executing specific tasks in space environments). Utilizing Mujoco (Todorov et al., 2012) for physics-based modeling, we systematically vary motion patterns and mechanical components to generate six datasets comprising 2200 trajectories, with each trajectory consisting of 120 time steps, enabling a thorough evaluation of the robustness of our approach. We describe the construction principles within Figure 5. Specifically, in the context of the Cubli robot, we have systematically varied the number of reaction wheels, creating distinct rigid body systems comprising single, double, and triple flywheels. For intricate rotational bodies, we have constructed three structures with no hinge, hinges on both sides, and hinges on the central position, respectively. All utilized physical systems are demonstrated within Appendix B.3

4.1.2. BASELINES

We conduct a comparative analysis of our proposed MDGS against various baselines, including GNS (Sanchez-Gonzalez et al., 2020), SGNN (Han et al., 2022a), LGNS (Bhattoo et al., 2022), and HGNS (Bishnoi et al., 2023). Specifically, GNS employs a pure Graph Neural Network (GNN) for simulating physical systems, while SGNN introduces the concept of subequivariant in the physical system into the GNN-based physical dynamics simulator. Furthermore, LGNS and HGNS leverage Lagrangian equations and Hamiltonian mechanics, respectively, to guide and constrain the GNN model, enhancing its capability to simulate physical reality. Apart from GNS, the other three methods incorporate varying degrees of prior knowledge from physics to reinforce the models. Through comparisons with these baselines, we can effectively analyze the algorithmic performance of our approach in the dynamic analysis of complex mechanical bodies.

4.1.3. EXPERIMENTAL SETTINGS

Due to the widely applicable enhancement provided by our designed rigid body structural constraint module on the accuracy of the GNN-based simulator in predicting rigid body problems, to ensure fairness in comparisons, we incorporated this module into other baselines as well, elevating their

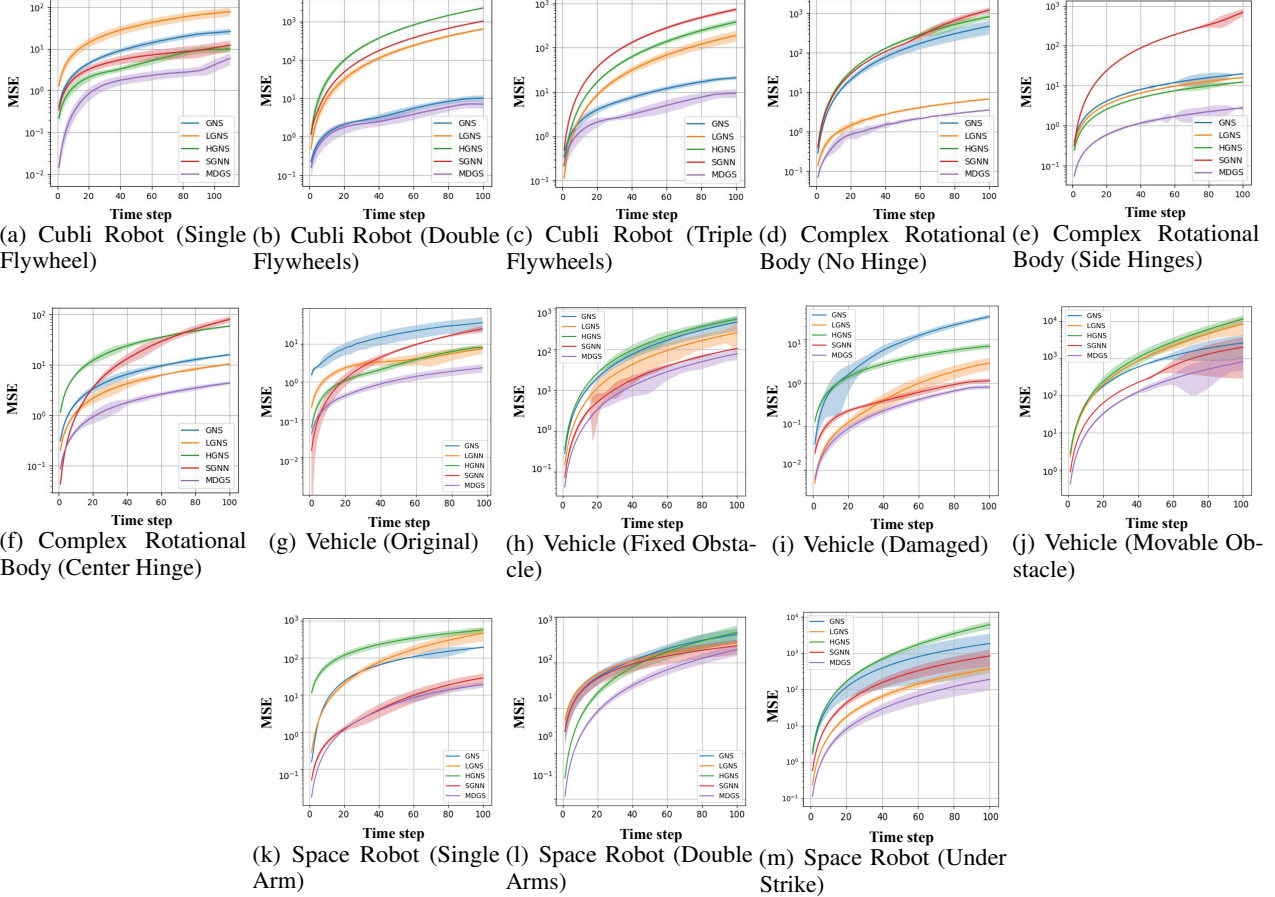


Figure 6. Experimental results on precision evaluation.

performance equivalently. Simultaneously, in Section 4.3.1, we presented the impact of adding or removing the rigid body structural constraint module on other baselines. Our experiments utilized MSE between predicted and ground truth values as the metric for evaluating the performance of each model. To demonstrate statistical significance, we followed the methodology outlined in (Han et al., 2022a), conducting each experiment 10 times and averaging the results. Specific hyperparameters and the backbones are detailed in Appendix B.2.

4.2. Performance Evaluation

4.2.1. PRECISION

Building upon the aforementioned experimental setup, we conducted performance testing experiments for both MDGS and other baseline algorithms across a total of six datasets. Figure 6 presents the experimental results. It is evident from the results that our proposed MDGS consistently outperforms the majority of comparisons. Figure 8 demonstrated the visualized results. We provide results represented in digital form in the Appendix B.4.

4.2.2. GENERALIZATION

We also conducted comparisons on the generalization performance of various methods. Using the complete Cubli Robot with three flywheels, we introduced variations to different attributes of the system. Subsequently, we examined whether the methods trained on the original Cubli Robot dataset could accurately predict the changed system and observed variations in algorithm prediction errors on the new test set. The test results are presented in the Figure ???. It is evident that our method achieves the best performance.

4.3. In-Depth Study

4.3.1. STRUCTURAL CONSTRAINT MODULE ANALYSIS

Validation experiments were conducted to analyze the capabilities of the proposed structural constraint module. A comparative study was performed on the dataset of the Cubli robot with all three flywheels, evaluating the performance of methods with and without the structural constraint module. In Figure 9, the performance of the respective models at different time steps is illustrated. It can be observed that all methods exhibit improved performance, i.e., lower MSE,

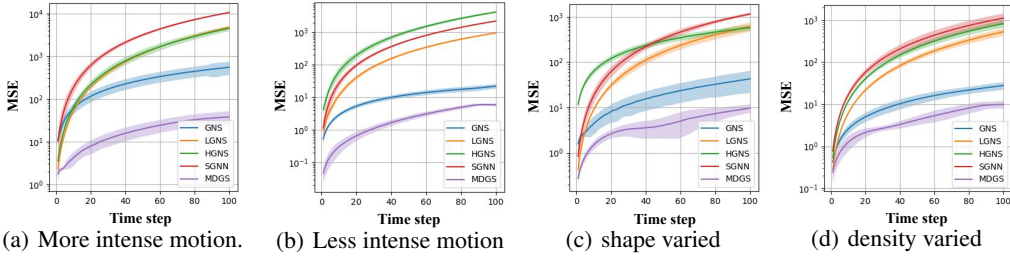


Figure 7. Experimental results on generalization.

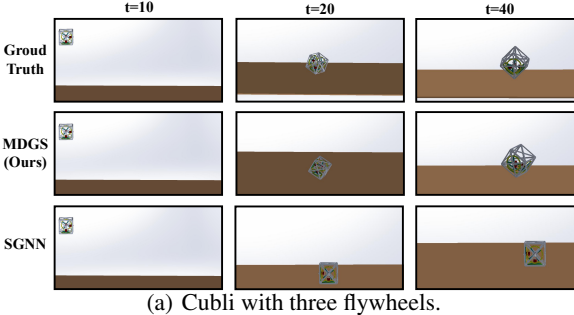


Figure 8. Visualized experimental results.

after incorporating the structural constraint module. This experimental result demonstrates the effectiveness of the proposed structural constraint module in enhancing the accuracy of model predictions in rigid body dynamics analysis, providing additional constraint information about the model structure for further predictions. Simultaneously, the experiment validates the necessity of integrating the structural constraint module when comparing our approach with other baselines. We provide results represented in digital form in the [Appendix B.4](#).

4.3.2. ABLATION STUDY

For a more in-depth analysis of our method, we conducted a series of ablation experiments. Specifically, we modified the constraints of Kane's equations to create two baseline models, namely MDGS-NC and MDGS-SC. MDGS-NC removes all constraints from Kane's equations, while MDGS-SC retains only an independent velocity constraint. The experimental results are depicted in Figure 10. It is evident that the constraints of Kane's equations indeed contribute to the improved performance of our method. Table 3 further illustrates the comparison between MDGS-NC, MDGS-SC,

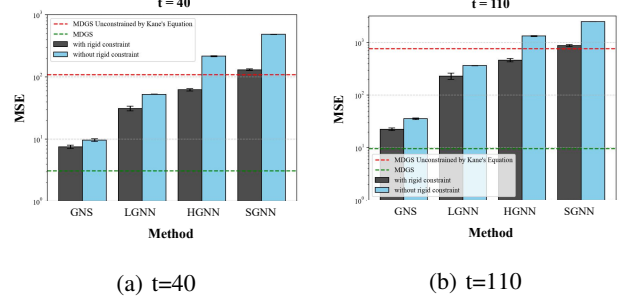


Figure 9. Experimental results of baselines with and without structural constraint module.

and other baselines.

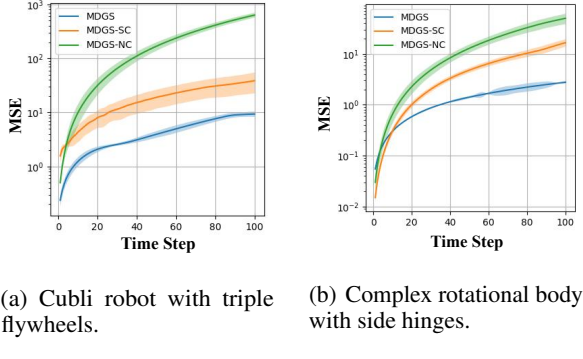


Figure 10. Ablation experiments results.

5. Conclusion

In this paper, we propose a novel approach, MDGS, to incorporate force analysis into GNN-based rigid body system simulators, aiming to enhance the accuracy of such methods in complex mechanical scenarios. We substantiate the design of MDGS through theoretical foundations and proofs and validate its performance through a series of experiments. We contend that MDGS can provide assistance to some extent in the application of GNN-based rigid body system simulators in industrial settings.

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Impact Statements

This paper presents work whose goal is to advance the field of Machine Learning. There are many potential societal consequences of our work, none of which we feel must be specifically highlighted here.

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A. Proofs

A.1. The Proof of Theorem 3.1

We first present the following lemma to assist in establishing the rationality of the theorem.

Lemma A.1. *By partitioning each rigid body within system \mathcal{S} into an element set $C = \{c_i\}_{i=0}^n$ following the procedure in Section 3.2.1, the motion states of all constructed elements within C , as well as the external forces $\{\vec{F}_i\}_{i=0}^n$ acting on these elements, conform to Kane's equation.*

Proof. According to the definition of generalized active force (Kane & Levinson, 1985), \mathcal{S} 's generalized active force K_γ according to the γ -th independent velocity can be expressed as follows:

$$K_\gamma = \sum_{k=1}^m \vec{F}_k \cdot \vec{u}_{k,\gamma}, \quad (16)$$

where m represents the number of point masses existing within \mathcal{S} , \vec{F}_k denotes the external force vector acting on point mass k , $\vec{u}_{k,\gamma}$ denotes the partial speed corresponding to the γ -th independent speed. The i -th element can be regarded as a rigid that consists of multiple point masses. Therefore, we have:

$$\begin{aligned} K_\gamma &= \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} \vec{F}'_{i,l} \cdot (\vec{u}_{i,\gamma} + \vec{\omega}_{i,\gamma} \times \vec{r}_{i,l}) = \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} \vec{F}'_{i,l} \cdot \vec{u}_{i,\gamma} + \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} \vec{F}'_{i,l} \cdot (\vec{\omega}_{i,\gamma} \times \vec{r}_{i,l}) \\ &= \sum_{i=1}^n \left(\sum_{l=1}^{\tilde{m}_i} \vec{F}'_{i,l} \right) \cdot \vec{u}_{i,\gamma} + \sum_{i=1}^n \left(\sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times \vec{F}'_{i,l}) \right) \cdot \vec{\omega}_{i,\gamma} = \sum_{i=1}^n \vec{R}_i \cdot \vec{u}_{i,\gamma} + \sum_{i=1}^n \vec{M}_i \cdot \vec{\omega}_{i,\gamma}, \end{aligned} \quad (17)$$

where $\vec{F}'_{i,l}$ denote the external force acting on l -th point mass among the \tilde{m}_i point masses that consist element c_i , $\vec{u}_{i,\gamma}$ denote the partial velocity according to the γ -th independent velocity acting on the mass center of the element c_i , $\vec{\omega}_{i,\gamma}$ denote the partial angular velocity of element c_i according to the γ -th independent velocity, $\vec{r}_{i,l}$ denotes the radius vector of the l -th point mass towards the mass center of the element c_i . \vec{R}_i and \vec{M}_i represent the forces and moments acting on individual elements. Such forces and moments can be calculated based on the properties of external forces $\{\vec{F}_i\}_{i=0}^n$. Consequently, we can derive that K_γ can be expressed as aggregating the external forces and moments acting on these elements.

Furthermore, we can represent \mathcal{S} 's generalized inertia force as follows:

$$K_\gamma^* = \sum_{k=1}^m (-m_k \vec{a}_k) \cdot \vec{u}_{k,\gamma}, \quad (18)$$

where m_k denote the mass of the k -th point mass, \vec{a}'_k represents its acceleration. Equation 18 can be reformulated as:

$$\begin{aligned} K_\gamma^* &= \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (-m_{i,l} \vec{a}_{i,l}) \cdot (\vec{u}_{i,\gamma} + \vec{\omega}_{i,\gamma} \times \vec{r}_{i,l}) = \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (-m_{i,l} \vec{a}_{i,l}) \cdot \vec{u}_{i,\gamma} + \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (-\vec{r}_{i,l} \times m_{i,l} \vec{a}_{i,l}) \cdot \vec{\omega}_{i,\gamma} \\ &= -\sum_{i=1}^n M_i \vec{a}_i \cdot \vec{u}_{i,\gamma} - \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times m_{i,l} \vec{a}_{i,l}) \cdot \vec{\omega}_{i,\gamma} = -\sum_{i=1}^n M_i \vec{a}_i \cdot \vec{u}_{i,\gamma} - \sum_{i=1}^n \vec{L}_i^* \cdot \vec{\omega}_{i,\gamma}, \end{aligned} \quad (19)$$

where $m_{i,l}$ denotes the mass of l -th point mass that consists element c_i , $\vec{a}_{i,l}$ denotes the corresponding acceleration. M_i and \vec{a}_i denotes the mass and the acceleration of c_i , \vec{L}_i^* can be determined based on the rotational state of c_i . Therefore, we can derive that K_γ^* can be expressed as the aggregation of the motion states of elements within C . As demonstrated afore, K_γ can be expressed as aggregating the external forces and moments acting on C . As both K_γ^* and K_γ within Kane's equation can be calculated with the given factors, we can conclude that the lemma holds. \square

Lemma A.1 demonstrates that the motion of element set C and external forces $\{\vec{F}_i\}_{i=0}^n$ conform to the Kane's equation. Next, we need to prove that equation 11 holds. According to the theorem, model $g(\cdot)$ is strictly accurate. Furthermore, from

605 Equation 17, we have:

$$606 \quad 607 \quad 608 \quad 609 \quad 610 \quad 611 \quad 612 \quad 613 \quad 614 \quad 615 \quad 616 \quad 617 \quad 618 \quad 619 \quad 620 \quad 621 \quad 622 \quad 623 \quad 624 \quad 625 \quad 626 \quad 627 \quad 628 \quad 629 \quad 630 \quad 631 \quad 632 \quad 633 \quad 634 \quad 635 \quad 636 \quad 637 \quad 638 \quad 639 \quad 640 \quad 641 \quad 642 \quad 643 \quad 644 \quad 645 \quad 646 \quad 647 \quad 648 \quad 649 \quad 650 \quad 651 \quad 652 \quad 653 \quad 654 \quad 655 \quad 656 \quad 657 \quad 658 \quad 659$$

$$\vec{R}_i = \sum_{l=1}^{\tilde{m}_i} \vec{F}_{i,l}, \quad (20)$$

which can be regarded as the resultant force acting on an element. Therefore, at time t , we have:

$$\vec{R}_i^{(t)} = [F_{1,i}^{(t)}, F_{2,i}^{(t)}, \dots, F_{d,i}^{(t)}]^\top, \quad (21)$$

where $F_{\gamma,i}^{(t)}$ denote the force value along coordinate q_γ at time t . Based on the statements in the theorem, due to the independence of generalized velocities from each other, we could take the generalized velocity corresponding to each generalized coordinate as independent velocity, the partial velocity $\vec{u}'_{i,\gamma}$ can be represented as follows:

$$\vec{u}'_{i,\gamma} = \frac{\partial \vec{r}_i}{\partial q_\gamma} = \frac{\partial [q_{1,i}, \dots, q_{d,i}]^\top}{\partial q_\gamma}, \quad (22)$$

where q_γ is the generalized coordinate corresponding to the γ -th independent velocity. $\vec{u}'_{i,\gamma}$ is a one-hot vector in such circumstances. Therefore, we have:

$$\vec{R}_i^{(t)} \cdot \vec{u}'_{i,\gamma} = [F_{1,i}^{(t)}, F_{2,i}^{(t)}, \dots, F_{d,i}^{(t)}]^\top \cdot \frac{\partial [q_{1,i}, \dots, q_{d,i}]^\top}{\partial q_\gamma} = F_{\gamma,i}^{(t)} u_\gamma, \quad (23)$$

where u_γ represents the unit velocity value along the direction of the γ -th independent velocity. Therefore, at time t , we have:

$$\begin{aligned} K_\gamma^{(t)} &= \sum_{i=1}^n \vec{r}_i \cdot \vec{u}'_{i,\gamma} + \sum_{i=1}^n \vec{M}_i^{(t)} \cdot \vec{\omega}_{i,\gamma}^{(t)} = \sum_{i=1}^n F_{\gamma,i}^{(t)} + \sum_{i=1}^n \vec{M}_i^{(t)} \cdot \vec{\omega}_{i,\gamma}^{(t)} \\ &= \sum_{i=1}^n F_{\gamma,i}^{(t)} + \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times \vec{F}_{i,l}^{(t)}) \cdot \vec{\omega}_{i,\gamma}^{(t)}. \end{aligned} \quad (24)$$

Similarly, we can also derive the following result:

$$K_\gamma^{*(t)} = - \sum_{i=1}^n M_i \vec{a}_i^{(t)} \cdot \vec{u}'_{i,\gamma} - \sum_{i=1}^n \vec{L}_i^{*(t)} \cdot \vec{\omega}_{i,\gamma}^{(t)} = - \sum_{i=1}^n M_i a_{\gamma,i}^{(t)} u_\gamma - \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times m_{i,l} \vec{a}_{i,l}^{(t)}) \cdot \vec{\omega}_{i,\gamma}^{(t)}, \quad (25)$$

where $a_{\gamma,i}^{(t)}$ denotes the accelerate speed value along coordinate q_γ at time t . According to Kane's equation, we have:

$$K_\gamma^{(t)} + K_\gamma^{*(t)} = 0. \quad (26)$$

With Equation 24, 25, and 26, we have:

$$\sum_{i=1}^n F_{\gamma,i}^{(t)} u_\gamma + \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times \vec{F}_{i,l}^{(t)}) \cdot \vec{\omega}_{i,\gamma}^{(t)} - \sum_{i=1}^n M_i a_{\gamma,i}^{(t)} u_\gamma - \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times m_{i,l} \vec{a}_{i,l}^{(t)}) \cdot \vec{\omega}_{i,\gamma}^{(t)} = 0. \quad (27)$$

As the prediction model $g(\cdot)$ has assumed to be able to accurately predict the state of \mathcal{S} in the Theorem 3.1, we have:

$$\sum_{i=1}^n F_{\gamma,i}^{(t)} u_\gamma - \sum_i M_i \frac{1}{\delta} (g(G^{(t)})_{[v_\gamma, i]} - v_\gamma^{(t)}) u_\gamma + \Psi^{(t)} = 0. \quad (28)$$

$\Psi^{(t)}$ is regarded as a disturbance term, formally:

$$\Psi^{(t)} = \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times \vec{F}_{i,l}^{(t)}) \cdot \vec{\omega}_{i,\gamma}^{(t)} - \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\vec{r}_{i,l} \times m_{i,l} \vec{a}_{i,l}^{(t)}) \cdot \vec{\omega}_{i,\gamma}^{(t)} \quad (29)$$

From Equation 29, it can be easily conclude that:

$$\Psi^{(t)} \propto \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} |\vec{r}_{i,l}|. \quad (30)$$

So far, we have demonstrated Equation 11, along with the property of $\Psi^{(t)}$.

660 A.2. The Proof of Theorem 3.2
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 662 Based on Equation 29, we have:
 663

$$\Psi^{(t)} = \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\mathbf{r}_{i,l} \times \vec{\mathbf{F}}_{i,l}^{(t)}) \cdot \boldsymbol{\omega}_{i,\gamma}^{(t)} - \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\mathbf{r}_{i,l} \times m_{i,l} \vec{\mathbf{a}}_{i,l}^{(t)}) \cdot \vec{\boldsymbol{\omega}}_{i,\gamma}^{(t)}. \quad (31)$$

 666 Within Equation 31, the first term of Ψ can be derived as follows:
 667

$$\begin{aligned} & \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\mathbf{r}_{i,l} \times \vec{\mathbf{F}}_{i,l}^{(t)}) \cdot \vec{\boldsymbol{\omega}}_{i,\gamma}^{(t)} \leq \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (|\mathbf{r}_{i,l}| |\vec{\mathbf{F}}_{i,l}^{(t)}| \sin(\theta_{i,l}^{RF})) \omega_\gamma \\ & \leq \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (|\mathbf{r}_{i,\arg \max_l(|\mathbf{r}_{i,l}|)}| |\vec{\mathbf{F}}_{i,l}^{(t)}| \sin(\theta_{i,l}^{RF})) \omega_\gamma = \sum_{i=1}^n |\mathbf{r}_{i,\arg \max_l(|\mathbf{r}_{i,l}|)}| \sum_{l=1}^{\tilde{m}_i} (|\vec{\mathbf{F}}_{i,l}^{(t)}| \sin(\theta_{i,l}^{RF})) \omega_\gamma \\ & \leq \mathbf{r}_{\arg \max_i |\mathbf{r}_{i,\arg \max_l(|\mathbf{r}_{i,l}|)}|, \arg \max_l(|\mathbf{r}_{i,l}|)} \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (|\vec{\mathbf{F}}_{i,l}^{(t)}| \sin(\theta_{i,l}^{RF})) \omega_\gamma, \end{aligned} \quad (32)$$

 678 where $\mathbf{r}_{\arg \max_i |\mathbf{r}_{i,\arg \max_l(|\mathbf{r}_{i,l}|)}|, \arg \max_l(|\mathbf{r}_{i,l}|)}$ is the max radius τ among all elements, $\theta_{i,l}^{RF}$ denotes the angle between $\mathbf{r}_{i,l}$
 679 and $\mathbf{F}_{i,l}^{(t)}$. The above expression can be further derived as follows:
 680

$$\tau \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (|\vec{\mathbf{F}}_{i,l}^{(t)}| \sin(\theta_{i,l}^{RF})) \omega_\gamma = \tau \sum_{i=1}^n (|\vec{\mathbf{R}}_i^{(t)}| \sin(\theta_i^{RF})) \leq \tau \sum_{i=1}^n (|\vec{\mathbf{R}}_i^{(t)}|) \omega_\gamma, \quad (33)$$

 684 where θ_i^{RF} denote the angle between $\vec{\mathbf{R}}_i^{(t)}$ and $\mathbf{r}_{i,l}$. Using the same method, we can also determine the lower bound of
 685 $\sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\mathbf{r}_{i,l} \times \vec{\mathbf{F}}_{i,l}^{(t)}) \cdot \vec{\boldsymbol{\omega}}_{i,\gamma}^{(t)}$, and thus we can obtain:
 686

$$-\tau \sum_{i=1}^n (|\vec{\mathbf{R}}_i^{(t)}|) \omega_\gamma \leq \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\mathbf{r}_{i,l} \times \vec{\mathbf{F}}_{i,l}^{(t)}) \cdot \vec{\boldsymbol{\omega}}_{i,\gamma}^{(t)} \leq \tau \sum_{i=1}^n (|\vec{\mathbf{R}}_i^{(t)}|) \omega_\gamma. \quad (34)$$

 691 Meanwhile, the second term of Equation 31 can be derived as follows:
 692

$$\sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\mathbf{r}_{i,l} \times m_{i,l} \vec{\mathbf{a}}_{i,l}^{(t)}) \cdot \vec{\boldsymbol{\omega}}_{i,\gamma}^{(t)} \leq \sum_{i=1}^n I_i \dot{\omega}_i^{(t)}, \quad (35)$$

 696 where I_i denotes the rotational inertia of element c_i . However, calculating the rotational inertia of irregular objects can be
 697 quite challenging, and this situation may arise with the elements we have divided. Therefore, we further derive Expressing
 698 35 as follows:
 699

$$\sum_{i=1}^n I_i \dot{\omega}_i^{(t)} \leq \sum_{i=1}^n \frac{2}{5} m_i \tau^2 \dot{\omega}_i^{(t)}, \quad (36)$$

 703 where $\frac{2}{5} m_i \tau^2$ be the rotational inertia of a sphere that contains element c_i . Based on Equation 35 and 36, we have:
 704

$$-\sum_{i=1}^n \frac{2}{5} m_i \tau^2 \dot{\omega}_i^{(t)} \leq \sum_{i=1}^n \sum_{l=1}^{\tilde{m}_i} (\mathbf{r}_{i,l} \times m_{i,l} \vec{\mathbf{a}}_{i,l}^{(t)}) \cdot \vec{\boldsymbol{\omega}}_{i,\gamma}^{(t)} \leq \sum_{i=1}^n \frac{2}{5} m_i \tau^2 \dot{\omega}_i^{(t)}, \quad (37)$$

 708 Therefore, we have:
 709

$$-\tau \sum_{i=1}^n (|\vec{\mathbf{R}}_i^{(t)}|) \omega_\gamma - \sum_{i=1}^n \frac{2}{5} m_i \tau^2 \dot{\omega}_i^{(t)} \leq \Psi \leq \tau \sum_{i=1}^n (|\vec{\mathbf{R}}_i^{(t)}|) \omega_\gamma + \sum_{i=1}^n \frac{2}{5} m_i \tau^2 \dot{\omega}_i^{(t)}, \quad (38)$$

 713 The theorem is proved.
 714

715 **B. Experimental Details**

716 **B.1. Implementations**

718 **B.1.1. EDGE CONSTRUCTION FOR CONTACT-ORIENTED GRAPH.**

719 In this section, we introduce the construction of edges in our CONTACT-ORIENTED GRAPH. Many physical scenarios are
 720 complex, involving multiple object interactions (including collision and friction) when the distance between them is minimal.
 721 The interactions between particles belonging to different objects and those within the same object are often distinct. Hence,
 722 we propose the concept of a contact graph. We continuously observe the distances between particles. When the distance
 723 between particles falls below a certain threshold, and they do not belong to the same object, we determine that contact has
 724 occurred. A new edge is established between these contacting particles to facilitate message passing, thus forming a new,
 725 contact-related graph. This newly constructed graph is utilized to predict and generate the contact forces experienced by
 726 the objects. Additionally, we leverage the pooling of particle-level information from the objects themselves to facilitate
 727 interactions between different objects.
 728

729 **B.1.2. THE SPECIFIC CALCULATION METHOD FOR EACH COMPONENT WITHIN \mathcal{L}_{cst} .**

730 For the prediction of $F_{\gamma,i}^{(t)}$, it is essential to first clarify the number of independent velocities in our system. The rigid body
 731 system under study is a six-degree-of-freedom system, encompassing x, y, and z coordinates, as well as angular velocities
 732 along the x, y, and z axes. Nevertheless, given the complexity of acquiring rotational inertia related to angular velocity and
 733 the challenging nature of obtaining torque data, we limit our calculations to determining the magnitudes of force components
 734 associated with independent velocities along the x, y, and z directions. These force components can be directly obtained
 735 from the corresponding elements of the force vector f_i calculated in Section 3.2.2. In this way, $|\vec{R}_i^{(t)}|$ can also be directly
 736 obtained by calculating the magnitude of the force vector f_i . M_i can be obtained by calculating the volume of each element
 737 and multiplying it by the density.
 738

739 We utilize the most commonly known laws of physics for solving angles issues. The angular velocity is derived by
 740 dividing the velocity difference of the current particle relative to the central particle by the distance between the current
 741 and the central particle. The formula is thus expressed as $\omega = \frac{\vec{v}}{r}$, where $\vec{v} = v_i(\text{vec, present}) - v_0(\text{vec, center}) =$
 742 $\sqrt{(v_{i,x} - v_{0,x})^2 + (v_{i,y} - v_{0,y})^2 + (v_{i,z} - v_{0,z})^2}$, $r = \sqrt{(x_i - x_0)^2 + (y_i - y_0)^2 + (z_i - z_0)^2}$. Similarly, for the
 743 velocity component of each axis, $\omega_{x,y,z} = \frac{v_{x,y,z}}{r_{x,y,z}}$, where $\vec{v}_x = v_{i,x} - v_{0,x}$, $\vec{v}_y = v_{i,y} - v_{0,y}$, $\vec{v}_z = v_{i,z} - v_{0,z}$;
 744 $r_x = x_i - x_0$, $r_y = y_i - y_0$, $r_z = z_i - z_0$. We will select the angular velocity component along the largest axis to
 745 use as our angular velocity, $\omega_{max} = \text{Max}(\omega_x, \omega_y, \omega_z)$.
 746

747 **B.1.3. STRUCTURAL CONSTRAINT MODULE**

748 In this study, we introduce a novel algorithm aimed at restoring deformed objects to their original, regular rigid body
 749 shapes. This algorithm is based on the geometric transformation of three-dimensional point sets, achieving the restoration
 750 of the object's shape by precisely calculating the optimal rotation and translation matrices. Specifically, the algorithm
 751 selects elements from the current and reference states of the object for processing. It calculates the centroid of each set
 752 and centralizes the data, shifting the coordinates of both sets to be centered around the origin. This step eliminates the
 753 influence of positional deviations. Subsequently, it computes the covariance matrix of the two centralized data sets and
 754 employs Singular Value Decomposition (SVD) to extract the rotation matrix. To ensure the correct rotational direction of the
 755 object, we introduce a correction matrix to adjust the rotation matrix. This rotation matrix not only reflects the directional
 756 differences between the two point sets but also ensures that the transformation adheres to the right-hand rule. Finally, by
 757 calculating the translation vector, the adjusted reference point set is moved to a position as close as possible to the current
 758 point set. Our experimental results indicate that this method can effectively restore deformed objects back to their original
 759 shapes.
 760

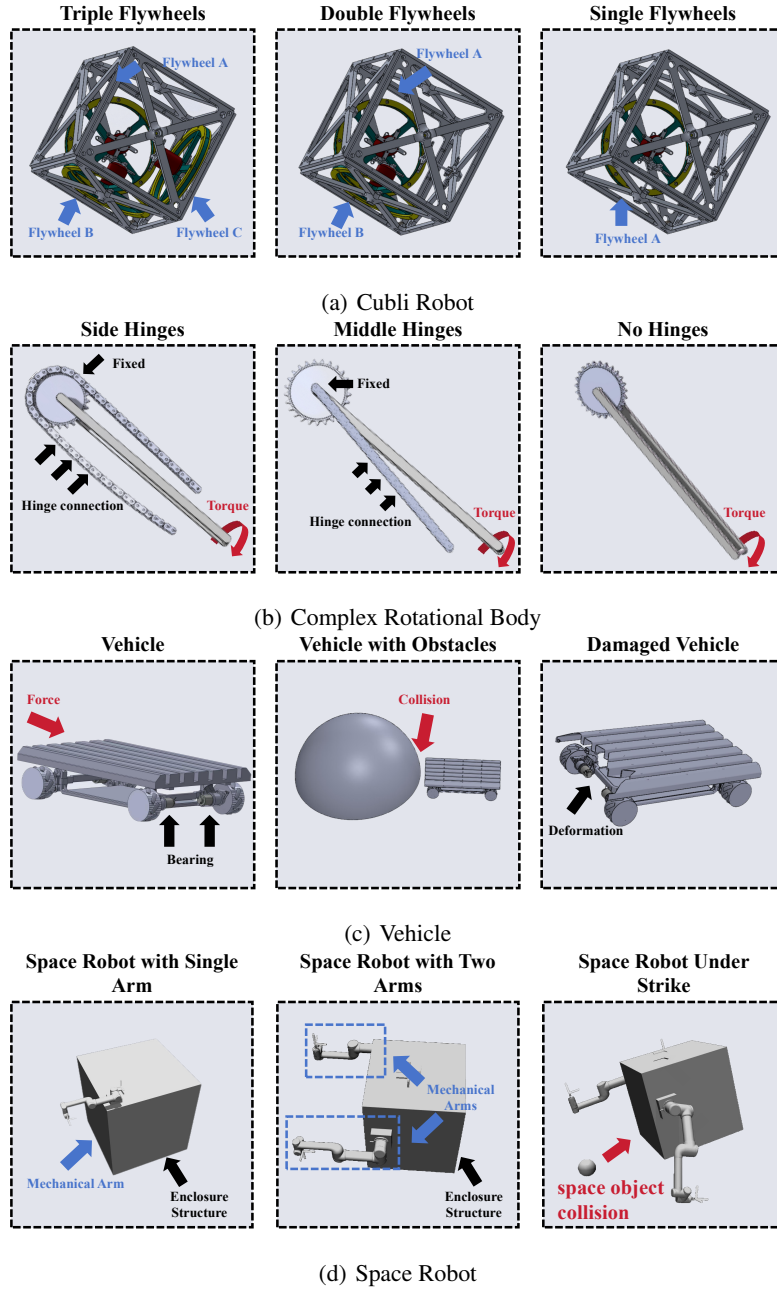
761 **B.2. Hyperparameters and Settings**

762 We adopt GCN as the backbone for all methods. Specifically, for LGNS and HGNS, we employ a hidden dimension of
 763 200 for the node update function and 300 for message computation, with each hidden dimension comprising 3 layers and
 764 utilizing Squareplus as the activation function. Due to its multi-stage hierarchical modeling, we use an Adam optimizer with
 765 an initial learning rate of 0.001 and beta values of (0.9, 0.999), along with a patience-based Plateau scheduler having 3 cycles
 766
 767

770 and a decay factor of 0.8. However, we still make very minor modifications based on the aforementioned hyperparameters.
 771 For SGNN, GNS, and MDGS, we employ SiLu or ReLU as the activation function and uniformly use a hidden dimension of
 772 200, with early stopping after 10 epochs. Given the larger size of each system, we use a batch size of 1 for each dataset.
 773 Furthermore, we inject noise during training to achieve better long-term prediction at test time, with the noise ratio set to
 774 4e-4 in Cubli, and 0.06 times the standard deviation in the other dataset. The cut-off radius r for both datasets is set to 0.001.
 775 On these two datasets, we only use the state information of the last frame t as input to predict the information of frame $t + 1$.
 776 Experiments are conducted on a single NVIDIA RTX A6000 GPU.
 777

778 B.3. Physical Systems Used in The Experiments

780 Figure 11 shows the detailed structures of all physical systems utilized in the experiments.
 781



823 *Figure 11.* The physical systems used in all twelve datasets.
 824

B.4. Further Results

Tables 1, 2, and 3 present the digital representation of the conducted experiments, to more clearly display the results.

Table 1. Comparative experiment results. **Bold** indicates the method with optimal performance. Underline denotes the method with second-best performance. Some standard deviations are marked as 0.00 due to being too small to effectively represent.

Scenario		Cubli Robot			Complex Rotational Body		
Time	Methods	Single Flywheel	Double Flywheels	Triple Flywheels	No Hinge	Side Hinges	Center Hinge
$t=20$ (MSE $\times 10^3$)	GNS	<u>4.47</u> ± 0.39	<u>2.08</u> ± 0.17	<u>3.95</u> ± 0.74	20.81 ± 3.14	4.09 ± 0.82	3.31 ± 0.96
	LGNS	14.05 ± 0.43	31.30 ± 6.80	8.52 ± 1.05	<u>1.42</u> ± 0.63	3.33 ± 0.86	<u>2.16</u> ± 1.15
	SGNN	3.22 ± 0.39	47.76 ± 3.55	36.27 ± 2.76	29.23 ± 5.29	23.86 ± 0.38	3.49 ± 0.47
	HGNS	<u>2.08</u> ± 0.69	97.02 ± 7.14	16.67 ± 0.81	33.61 ± 6.33	<u>2.55</u> ± 0.37	12.18 ± 2.05
	MDGS (Ours)	0.84 ± 0.28	1.76 ± 0.63	2.08 ± 0.24	0.84 ± 0.04	0.58 ± 0.03	0.92 ± 0.15
$t=40$ (MSE $\times 10^3$)	GNS	0.89 ± 0.17	<u>0.32</u> ± 0.26	<u>0.75</u> ± 0.05	7.77 ± 1.79	0.80 ± 0.09	0.65 ± 0.19
	LGNS	2.80 ± 1.08	11.00 ± 1.73	3.12 ± 0.26	<u>0.27</u> ± 0.01	0.65 ± 0.22	<u>0.42</u> ± 0.22
	SGNN	0.54 ± 0.32	17.24 ± 1.12	13.09 ± 0.38	10.65 ± 0.21	8.73 ± 1.06	1.31 ± 0.83
	HGNS	<u>0.33</u> ± 0.13	36.51 ± 2.29	6.24 ± 0.27	12.84 ± 0.52	<u>0.50</u> ± 0.09	2.37 ± 0.98
	MDGS (Ours)	0.17 ± 0.08	0.24 ± 0.06	0.31 ± 0.03	0.15 ± 0.04	0.11 ± 0.08	0.18 ± 0.04
$t=110$ (MSE $\times 10^2$)	GNS	0.25 ± 0.11	<u>0.14</u> ± 0.02	<u>0.22</u> ± 0.01	5.71 ± 2.37	0.22 ± 0.04	0.17 ± 0.03
	LGNS	0.77 ± 0.23	7.61 ± 0.92	2.30 ± 1.13	<u>0.07</u> ± 0.01	0.18 ± 0.03	<u>0.11</u> ± 0.02
	SGNN	0.12 ± 0.07	12.20 ± 0.37	8.73 ± 0.79	15.27 ± 1.21	9.54 ± 1.06	0.96 ± 0.21
	HGNS	<u>0.09</u> ± 0.01	26.79 ± 0.44	4.61 ± 0.35	10.13 ± 0.78	<u>0.14</u> ± 0.01	0.64 ± 0.39
	MDGS (Ours)	0.05 ± 0.01	0.06 ± 0.01	0.09 ± 0.01	0.04 ± 0.01	0.03 ± 0.00	0.05 ± 0.00

Table 2. Generalization experiment results. Δ indicates the disparity in model performance between the test set with changes and the original test set. **Bold** indicates the method with optimal performance. Underline denotes the method with second-best performance.

Time	Methods	More Intense Motion	Δ	Less Intense Motion	Δ	Shape Varied	Δ	Density Varied	Δ
$t=20$ (MSE $\times 10^3$)	GNS	<u>115.61</u> ± 18.08	<u>+113.53</u>	5.41 ± 0.27	<u>+3.33</u>	<u>5.01</u> ± 0.62	<u>+1.06</u>	<u>8.17</u> ± 3.57	<u>+4.22</u>
	LGNS	188.71 ± 28.27	<u>+157.41</u>	40.79 ± 13.93	<u>+9.49</u>	21.77 ± 0.83	<u>+13.25</u>	30.72 ± 9.43	<u>+22.20</u>
	SGNN	614.13 ± 31.04	566.36	97.02 ± 49.25	<u>+49.26</u>	57.53 ± 1.02	<u>+21.26</u>	61.92 ± 48.41	<u>+25.65</u>
	HGNS	216.76 ± 47.45	<u>+119.74</u>	197.58 ± 61.12	<u>+100.56</u>	40.53 ± 1.02	<u>+23.86</u>	120.01 ± 34.54	<u>+103.34</u>
	MDGS (Ours)	7.91 ± 2.85	+6.14	4.47 ± 0.38	+2.71	2.13 ± 0.26	+0.05	2.63 ± 0.58	+0.55
$t=40$ (MSE $\times 10^3$)	GNS	<u>22.51</u> ± 3.15	<u>+22.19</u>	<u>1.00</u> ± 0.15	<u>+0.68</u>	1.03 ± 0.13	<u>+0.28</u>	<u>1.55</u> ± 0.72	<u>+0.79</u>
	LGNS	73.64 ± 2.51	<u>+62.63</u>	15.62 ± 3.44	<u>+4.61</u>	8.22 ± 1.21	<u>+5.10</u>	11.05 ± 1.98	<u>+7.93</u>
	SGNN	211.42 ± 19.05	<u>+194.18</u>	36.51 ± 6.29	<u>+19.26</u>	20.61 ± 3.53	<u>+7.52</u>	22.02 ± 3.08	<u>+8.93</u>
	HGNS	78.16 ± 16.71	<u>+41.66</u>	70.53 ± 21.02	<u>+33.54</u>	14.68 ± 2.55	<u>+8.44</u>	23.37 ± 2.88	<u>+17.13</u>
	MDGS (Ours)	1.57 ± 0.56	+1.32	0.90 ± 0.19	+0.64	0.33 ± 0.03	+0.02	0.36 ± 0.15	+0.05
$t=110$ (MSE $\times 10^2$)	GNS	6.08 ± 0.05	<u>+5.97</u>	<u>0.36</u> ± 0.14	<u>+0.22</u>	<u>0.31</u> ± 0.03	<u>+0.09</u>	<u>0.48</u> ± 0.26	<u>+0.26</u>
	LGNS	58.33 ± 4.91	<u>+50.72</u>	11.60 ± 0.15	<u>+3.97</u>	6.41 ± 1.32	<u>+4.11</u>	7.25 ± 0.16	<u>+4.95</u>
	SGNN	124.04 ± 11.04	<u>+111.83</u>	26.80 ± 2.47	<u>+1.45</u>	13.11 ± 2.37	<u>+4.38</u>	13.67 ± 0.02	<u>+4.94</u>
	HGNS	54.40 ± 7.15	<u>+28.61</u>	50.86 ± 8.32	<u>+24.07</u>	9.76 ± 0.98	<u>+5.15</u>	6.37 ± 0.24	<u>+1.76</u>
	MDGS (Ours)	0.40 ± 0.15	+0.33	0.25 ± 0.02	+0.19	0.10 ± 0.01	+0.01	0.11 ± 0.02	+0.02

Table 3. Ablation experiments compared with other baselines.

Methods	Cubli Robot		Complex Rotational Body	
	t=40	t=110	t=40	t=110
GNS	7.52 ± 0.49	22.36 ± 1.21	8.01 ± 0.69	21.70 ± 3.71
LGNS	31.2 ± 2.66	230.09 ± 31.94	6.52 ± 0.87	17.59 ± 2.52
SGNN	130.95 ± 3.77	873.55 ± 39.42	87.38 ± 7.26	954.46 ± 60.82
HGNS	62.44 ± 2.68	461.5 ± 35.41	4.98 ± 0.19	13.53 ± 1.07
MDGS-NC	110.84 ± 12.47	763.36 ± 1.46	8.40 ± 0.33	60.15 ± 7.05
MDGS-SC	15.24 ± 5.88	43.47 ± 18.94	3.32 ± 0.36	20.85 ± 3.27
MDGS	3.11 ± 0.32	9.59 ± 1.27	1.15 ± 0.31	3.04 ± 0.64