K. Scalability with respect to Dimension and **Constraints**

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We first present the results of exact sampling, computing closed-form loss, and gradient estimator with respect to varying dimensions. The number of constraints is half of the dimension (n). All results are averaged over 100 runs and conducted on a Nvidia A800 80GB GPU. All numbers are measured in seconds. We highlight that even though increasing dimension slows down the algorithm, the additional computational cost can be mitigated by choosing a larger batch size. For example, the additional time taken to run with batch_size=16 and batch_size=32 is almost negligible for n=32,64, and 128. In the challenging n = 2048 case, using batch_size=32 only takes 1.5 times more time to run compared to batch size=16. Larger batch sizes can significantly reduce computation time per sample thanks to the parallelization of our method.

	16	32	64	128	256
n = 32	0.00249	0.00250	0.00252	0.00253	0.00255
n = 64	0.00348	0.00348	0.00348	0.00352	0.00366
n = 128	0.00467	0.00473	0.00479	0.00483	0.00498
n = 256	0.00760	0.00769	0.00806	0.00916	0.01052
n = 512	0.00760	0.01770	0.02159	0.02921	0.04500
n = 1024	0.04843	0.06266	0.09138	0.13943	0.23699
n = 2048	0.20515	0.31654	0.46224	0.78970	1.24126

Table 11. Transposed computation time (in seconds) for varying dimensions and batch sizes.

```
Algorithm 2 Varying Dimensions
```

13: **end for**

```
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        1: for each n \in ls\_n do
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              num\_constraints \leftarrow n/2
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              Total\ time \leftarrow 0
        3:
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        4:
              for i \leftarrow 1 to repeat\_num do
1135
                 Generate unconstrained parameters and constraints
        5:
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                 Conduct exact sampling
        6:
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                 Compute closed-form loss
        7:
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                 Compute gradient estimator
        8:
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                 Total \ time \leftarrow Total \ time + time
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       10:
              end for
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       11:
              Total\_time \leftarrow Total\_time/repeat\_num
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              print(Total time)
       12:
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```

We next investigate the impact of the number of constraints (num k). We conduct experiments on the challenging n=2048 setting. Similar to the findings in scaling the number of dimensions, using larger batch sizes significantly reduces computation time per sample. Additionally, we also found that increasing the number of constraints does not significantly increase the computational time. For example, under batch_size = 64, increasing the number of constraints from 256 to 512 only increases computational time by 32%.

	16	32	64	128	256
$\texttt{num_k} = 32$	0.05109	0.09899	0.19531	0.29093	0.47706
${\tt num_k}=64$	0.05343	0.10259	0.20241	0.30316	0.48106
$\texttt{num_k} = 128$	0.05894	0.11060	0.21666	0.32456	0.50057
${\tt num_k} = 256$	0.07336	0.13211	0.24871	0.38556	0.52001
$\texttt{num_k} = 512$	0.10885	0.18119	0.32935	0.52796	0.84375
$\texttt{num_k} = 1024$	0.20515	0.31654	0.46224	0.78970	1.24126

Table 12. Transposed computation time (in seconds) for varying number of constraints and batch sizes at n = 2048.

Algorithm 3 Exact Sampling and Gradient Estimation over Varying Number of Constraints

```
1: for each num\_k \in ls\_k do
      Total\ time \leftarrow 0
2:
3:
      for i \leftarrow 1 to repeat num do
         Generate unconstrained parameters and constraints
4:
5:
         Conduct exact sampling
         Compute closed-form loss
6:
7:
         Compute gradient estimator
         Total\_time \leftarrow Total\_time + time
8:
9:
      Total\_time \leftarrow Total\_time/repeat\_num
10:
      print(Total time)
11:
12: end for
```

L. Additional Experiment on CL and **Constrained Reparametrization**

Additional experiment results on CL and Constrained Reparametrization are highlighted with gray.

Table 13. Comparison on VAE generative performance. The constrained VAE models achieve similar or better generative ability while strictly satisfying the constraints, whereas the unconstrained counterparts have a high constraint violation rate.

MODEL	LL ↑	ELBO ↑	RL ↓	Violation ↓
VAE	-22.42 ± 0.29	-23.41 ± 0.22	15.00 ± 0.46	0.30 ± 0.06
VAE + Constr Layer	-34.45 ± 2.6	$4 -40.89 \pm 9.37$	37.11 ± 9.35	$\textbf{0.00} \pm \textbf{0.00}$
VAE + Constr Reparam	-22.33 ± 0.48	-23.83 ± 0.49	14.54 ± 0.58	0.00 ± 0.00
ours	$\textbf{-21.48} \pm \textbf{0.1}$	-22.62 ± 0.07	$\textbf{12.79} \pm \textbf{0.11}$	$\textbf{0.00} \pm \textbf{0.00}$
Ladder VAE	-24.25 ± 0.0°	$7 -30.84 \pm 0.51$	$\textbf{23.06} \pm \textbf{0.54}$	0.38 ± 0.02
Ladder VAE + Constr Layer	-36.83 ± 0.56	$6 -39.59 \pm 0.56$	37.46 ± 0.55	$\textbf{0.00} \pm \textbf{0.00}$
Ladder VAE + Constr Reparam	-25.27 ± 0.19	-31.56 ± 0.48	25.20 ± 0.62	0.00 ± 0.00
ours	-23.86 ± 0.0	$6 -30.78 \pm 0.08$	$\textbf{23.40} \pm \textbf{0.16}$	0.00 ± 0.00
Graph VAE	-22.74 ± 0.1	-23.54 ± 0.18	15.45 ± 0.41	0.29 ± 0.09
Graph VAE + Constr Layer	-33.27 ± 3.66	-33.27 ± 5.84	28.29 ± 6.40	$\textbf{0.00} \pm \textbf{0.00}$
Graph VAE + Constr Reparam	-22.96 ± 0.80	-23.55 ± 0.86	16.08 ± 1.38	0.00 ± 0.00
ours	-21.61 ± 0.2	-22.53 ± 0.06	$\textbf{12.73} \pm \textbf{0.21}$	$\textbf{0.00} \pm \textbf{0.00}$

L.1. Constrained Generation using VAE

L.3. Charge-Neutral Predictions

L.2. Constrained Generation using Diffusion Models

Table 14. Comparison of constrained and unconstrained models across datasets under DDPM. We report FID (Fréchet Inception Distance), IS (Inception Score), and Violation metrics.

DATASET	MODEL	FID \downarrow	IS ↑	$\textbf{Violation} \downarrow$
	Ours	3.811	9.223 ± 0.130	0
CIFAR	Constr Layer	4.234	8.535 ± 0.157	0
011.111	Constr Reparam	3.881	9.212 ± 0.122	0
	DDPM	4.173	$\textbf{9.278} \pm \textbf{0.116}$	0.999
	Ours	10.193	$\textbf{2.360} \pm \textbf{0.016}$	0
CelebA	Constr Layer	12.067	2.345 ± 0.039	0
CCICDA	Constr Reparam	10.961	2.043 ± 0.028	0
	DDPM	10.345	2.358 ± 0.030	0.999
	Ours	4.779	2.471 ± 0.020	0
LSUN Church	Constr Layer	6.695	2.319 ± 0.028	0
	Constr Reparam	4.895	2.377 ± 0.032	0
	DDPM	4.945	2.460 ± 0.028	1.0
	Ours	12.489	$\textbf{4.711} \pm \textbf{0.054}$	0
LSUN Cat	Constr Layer	13.472	4.642 ± 0.081	0
	Constr Reparam	12.696	4.707 ± 0.062	0
	DDPM	12.913	4.705 ± 0.047	1.0

Table 15. Performances of different methods for estimating partial charges on metal ions are presented. Compared to the baseline MPNN (variance), both the closed-form loss function and likelihood objective yield superior mean absolute deviation (MAD) results. The same holds for their ensemble counterpart. We find that ensemble methods (second block) notably boost the predictive performance in general.

METHOD neutrality enforcement	$\mathbf{MAD}\downarrow\\\mathbf{mean}\pm\mathbf{std}$	$\mathbf{NLL}\downarrow\\\mathbf{mean}\pm\mathbf{std}$
Constrained Layer	0.327 ± 0.004	103.522 ± 3.018
Constant Prediction	0.324 ± 0.007	_
Element-mean (uniform)	0.154 ± 0.002	_
Element-mean (variance)	0.153 ± 0.002	_
MPNN (KKThPINN)	0.0260 ± 0.0008	109.8 ± 6.9
MPNN (Constr Reparam)	0.0256 ± 0.0007	$3340.03 \;\pm\; 2398$
MPNN (variance)	0.0251 ± 0.0010	-19.9 ± 71.1
Closed-form (ours)	0.0245 ± 0.0009	> 1e+7
Likelihood (ours)	0.0248 ± 0.0008	-252 ± 24.7
Constrained Layer (ens)	0.319 ± 0.002	99.236 ± 2.3
MPNN (ens, KKThPINN)	0.0244 ± 0.0006	57.29 ± 12.8
MPNN (ens, Constr Reparam)	0.0242 ± 0.0006	4883.46 ± 1695
MPNN (ens, variance)	0.0238 ± 0.0007	-45.2 ± 55.8
Closed-form (ens, ours)	0.0230 ± 0.0008	>1e+7
Likelihood (ens, ours)	0.0231 ± 0.0007	-180 ± 38.3

1210 L.4. Chemical Process Units and Subsystems

Table 16. Comparison of models across **CSTR**, **plant**, and **distillation** tasks. The mean and standard deviation of MSE scaled by 10^{-4} are reported. All experiments are averaged for 10 times.

MODEL	CSTR	PLANT	DISTILLATION
ECNN	20.6 ± 27.0	0.31 ± 0.23	1.94 ± 0.70
KKThPINN	11.7 ± 20.3	0.11 ± 0.04	2.02 ± 0.94
NN	18.3 ± 20.8	0.34 ± 0.64	1.99 ± 0.67
PINN	260.8 ± 20.4	3.62 ± 1.94	40.9 ± 10.7
CL	9.28 ± 3.56	0.58 ± 0.64	2.26 ± 1.19
Constr Reparam	7.13 ± 3.22	$0.14\ \pm\ 0.07$	$2.22\ \pm\ 0.94$
Ours	$\textbf{4.31} \pm \textbf{1.58}$	$\textbf{0.09} \pm \textbf{0.05}$	$\textbf{1.73} \pm \textbf{0.70}$

L.5. Stock Investment

Table 17. Comparison of models based on Sharpe ratio. We report the mean and standard deviation averaged across 10 runs.

MODEL	SHARPE RATIO ↑		
StemGNN	1.5576 ± 0.3405		
StemGNN-KKThPINN	1.8092 ± 0.7055		
StemGNN-CL	1.5018 ± 0.3318		
StemGNN-Constr Reparam	1.8162 ± 0.2966		
Ours	1.9041 ± 0.2329		