GAN (Generative Adversarial Nets)

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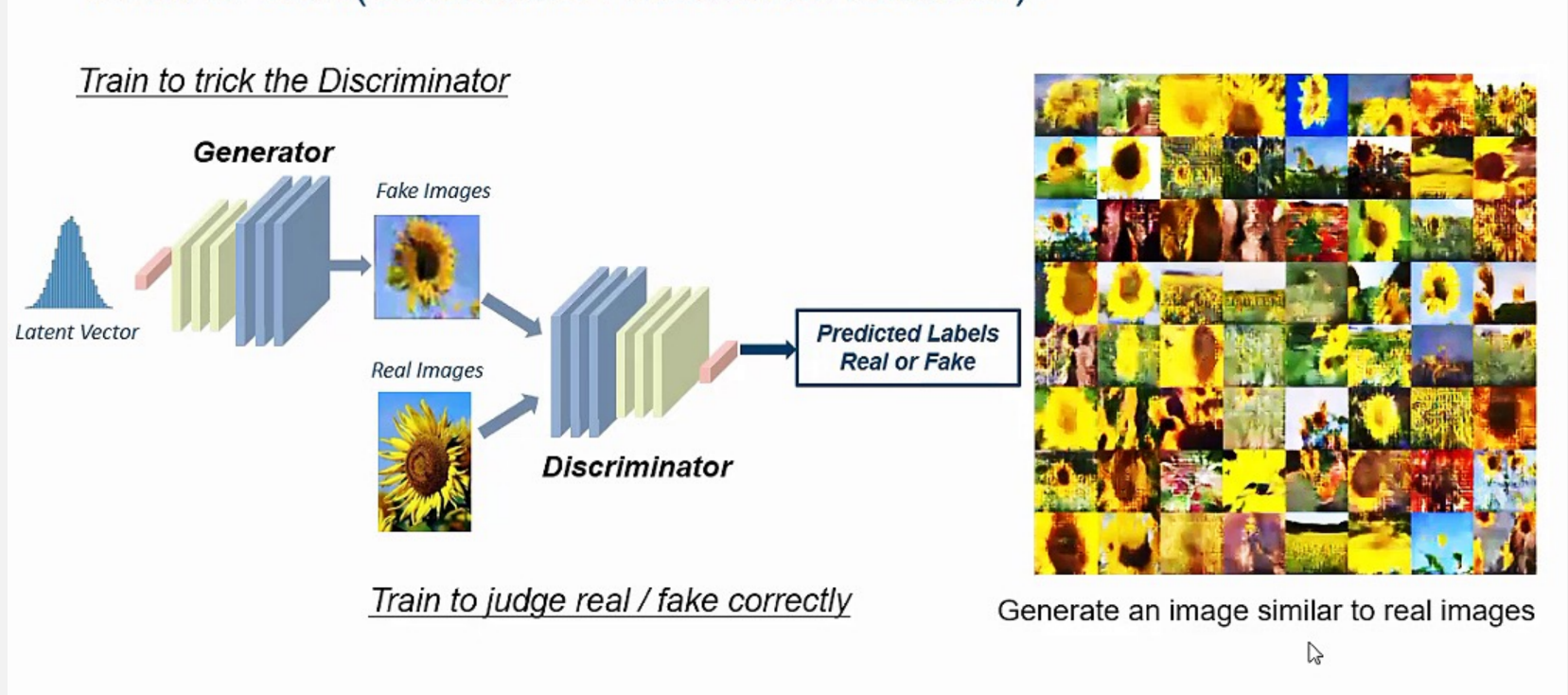
2021.07.12

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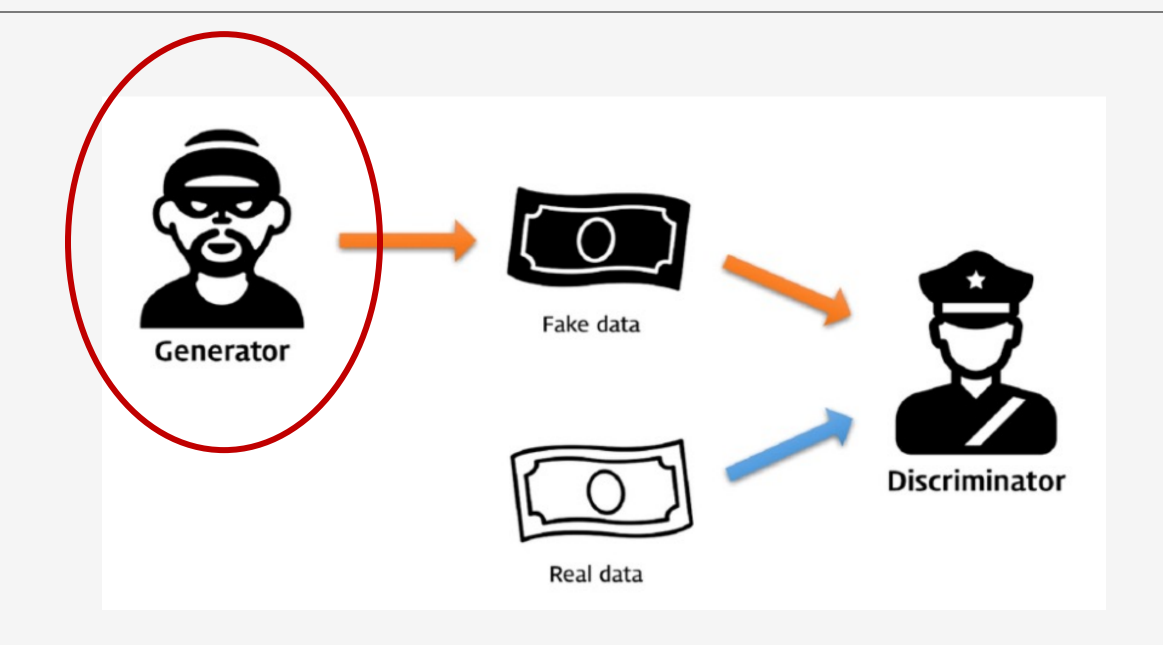
곽수지

GAN(Generative Adversarial Networks, 생성적 적대 신경망)

What is GAN(Generative Adversarial Network)?



Overview

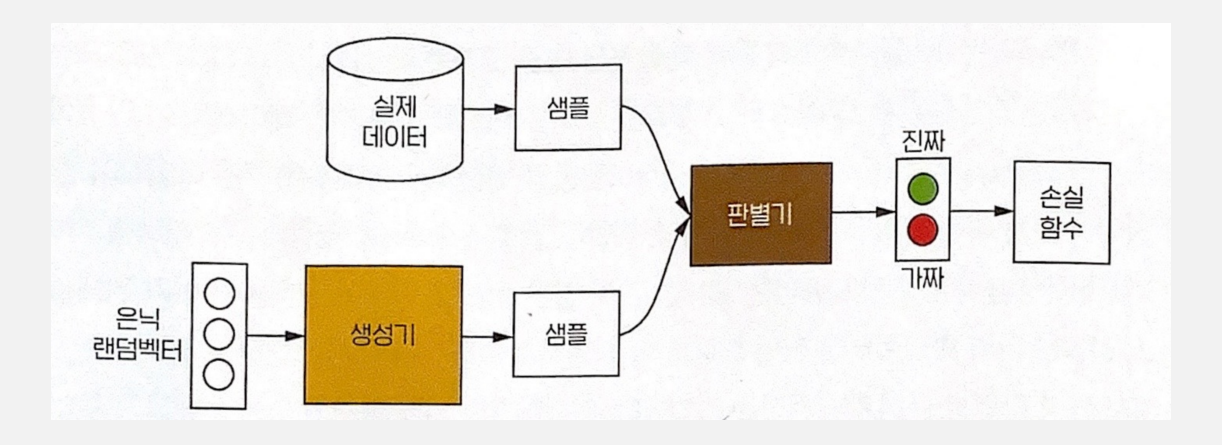


Keywords: MinMax 게임 MinMax Two-Player Game / 가치 함수 Value Function /

JS 거리 Jensen-Shannon Divergence / 확률생성모델 Probability Generating Model /

최적화 포화 Optimization Saturation

생성 모델과 판별 모델의 목적

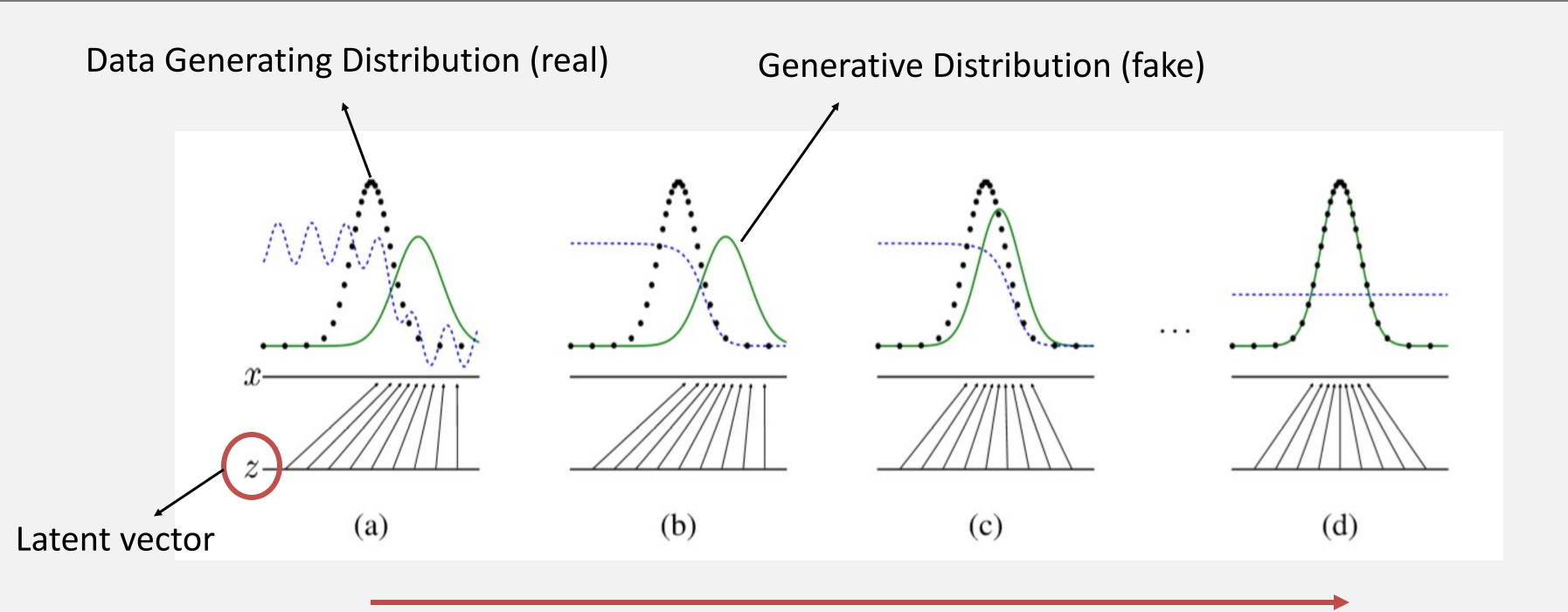


Generator(생성 모델):

생성한 데이터가 실제 데이터의 확률 분포와 같아지도록 학습

Discriminator(판별 모델):

샘플 데이터가 G로부터 나온 데이터가 아닌 실제 데이터로부터 나온 데이터일 확률을 추정



생성 모델 G가 학습 데이터의 분포를 학습함

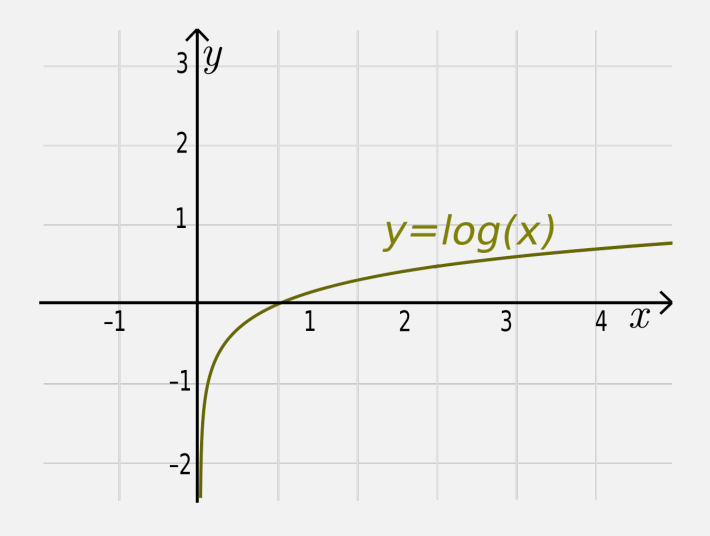
Value Function

$$\min_{G} \max_{D} V(D,G) = \mathbb{E}_{\boldsymbol{x} \sim p_{\text{data}}(\boldsymbol{x})}[\log D(\boldsymbol{x})] + \mathbb{E}_{\boldsymbol{z} \sim p_{\boldsymbol{z}}(\boldsymbol{z})}[\log (1 - D(G(\boldsymbol{z})))]$$

MinMax Two Player Game

D: value function 최대화 (→ 0)

G: value function 최소화 $(\rightarrow -\infty)$



Algorithm

Algorithm 1 Minibatch stochastic gradient descent training of generative adversarial nets. The number of steps to apply to the discriminator, k, is a hyperparameter. We used k=1, the least expensive option, in our experiments.

for number of training iterations do

for k steps do

- Sample minibatch of m noise samples $\{z^{(1)}, \dots, z^{(m)}\}$ from noise prior $p_g(z)$.
- Sample minibatch of m examples $\{x^{(1)}, \dots, x^{(m)}\}$ from data generating distribution $p_{\text{data}}(x)$.
- Update the discriminator by ascending its stochastic gradient:

$$\nabla_{\theta_d} \frac{1}{m} \sum_{i=1}^m \left[\log D\left(\boldsymbol{x}^{(i)} \right) + \log \left(1 - D\left(G\left(\boldsymbol{z}^{(i)} \right) \right) \right) \right].$$

end for

- Sample minibatch of m noise samples $\{z^{(1)}, \dots, z^{(m)}\}$ from noise prior $p_q(z)$.
- Update the generator by descending its stochastic gradient:

$$\nabla_{\theta_g} \frac{1}{m} \sum_{i=1}^{m} \log \left(1 - D \left(G \left(\boldsymbol{z}^{(i)} \right) \right) \right).$$

end for

The gradient-based updates can use any standard gradient-based learning rule. We used momentum in our experiments.

K 스텝마다 **D** 최적화 1 스텝마다 **G** 최적화

- → **D** overfitting 방지
- → Computational Problem

Theoretical Results

4.1 Global Optimality of $p_a = p_{data}$

We first consider the optimal discriminator D for any given generator G.

Proposition 1. For G fixed, the optimal discriminator D is

$$D_G^*(\mathbf{x}) = \frac{p_{data}(\mathbf{x})}{p_{data}(\mathbf{x}) + p_q(\mathbf{x})}$$
(2)

Proof. The training criterion for the discriminator D, given any generator G, is to maximize the quantity V(G,D)

$$V(G, D) = \int_{\mathbf{x}} p_{\text{data}}(\mathbf{x}) \log(D(\mathbf{x})) dx + \int_{z} p_{\mathbf{z}}(\mathbf{z}) \log(1 - D(g(\mathbf{z}))) dz$$
$$= \int_{\mathbf{x}} p_{\text{data}}(\mathbf{x}) \log(D(\mathbf{x})) + p_{g}(\mathbf{x}) \log(1 - D(\mathbf{x})) dx$$
(3)

For any $(a,b) \in \mathbb{R}^2 \setminus \{0,0\}$, the function $y \to a \log(y) + b \log(1-y)$ achieves its maximum in [0,1] at $\frac{a}{a+b}$. The discriminator does not need to be defined outside of $Supp(p_{\text{data}}) \cup Supp(p_g)$, concluding the proof.

Note that the training objective for D can be interpreted as maximizing the log-likelihood for estimating the conditional probability P(Y = y|x), where Y indicates whether x comes from p_{data} (with y = 1) or from p_g (with y = 0). The minimax game in Eq. 1 can now be reformulated as:

$$C(G) = \max_{D} V(G, D)$$

$$= \mathbb{E}_{\boldsymbol{x} \sim p_{\text{data}}} [\log D_{G}^{*}(\boldsymbol{x})] + \mathbb{E}_{\boldsymbol{z} \sim p_{\boldsymbol{z}}} [\log (1 - D_{G}^{*}(G(\boldsymbol{z})))]$$

$$= \mathbb{E}_{\boldsymbol{x} \sim p_{\text{data}}} [\log D_{G}^{*}(\boldsymbol{x})] + \mathbb{E}_{\boldsymbol{x} \sim p_{g}} [\log (1 - D_{G}^{*}(\boldsymbol{x}))]$$

$$= \mathbb{E}_{\boldsymbol{x} \sim p_{\text{data}}} \left[\log \frac{p_{\text{data}}(\boldsymbol{x})}{P_{\text{data}}(\boldsymbol{x}) + p_{g}(\boldsymbol{x})} \right] + \mathbb{E}_{\boldsymbol{x} \sim p_{g}} \left[\log \frac{p_{g}(\boldsymbol{x})}{p_{\text{data}}(\boldsymbol{x}) + p_{g}(\boldsymbol{x})} \right]$$
(4)

Global Optimality of $P_g = P_{data}$

Theorem 1. The global minimum of the virtual training criterion C(G) is achieved if and only if $p_a = p_{data}$. At that point, C(G) achieves the value $-\log 4$.

Proof. For $p_g = p_{\text{data}}$, $D_G^*(x) = \frac{1}{2}$, (consider Eq. 2). Hence, by inspecting Eq. 4 at $D_G^*(x) = \frac{1}{2}$, we find $C(G) = \log \frac{1}{2} + \log \frac{1}{2} = -\log 4$. To see that this is the best possible value of C(G), reached only for $p_g = p_{\text{data}}$, observe that

$$\mathbb{E}_{\boldsymbol{x} \sim p_{\text{data}}} \left[-\log 2 \right] + \mathbb{E}_{\boldsymbol{x} \sim p_{a}} \left[-\log 2 \right] = -\log 4$$

and that by subtracting this expression from $C(G) = V(D_G^*, G)$, we obtain:

$$C(G) = -\log(4) + KL\left(p_{\text{data}} \left\| \frac{p_{\text{data}} + p_g}{2} \right) + KL\left(p_g \left\| \frac{p_{\text{data}} + p_g}{2} \right) \right)$$
 (5)

where KL is the Kullback–Leibler divergence. We recognize in the previous expression the Jensen–Shannon divergence between the model's distribution and the data generating process:

$$C(G) = -\log(4) + 2 \cdot JSD\left(p_{\text{data}} \| p_{\alpha}\right) \tag{6}$$

Since the Jensen–Shannon divergence between two distributions is always non-negative and zero only when they are equal, we have shown that $C^* = -\log(4)$ is the global minimum of C(G) and that the only solution is $p_g = p_{\text{data}}$, i.e., the generative model perfectly replicating the data generating process.

4.2 Convergence of Algorithm 1

Proposition 2. If G and D have enough capacity, and at each step of Algorithm 1, the discriminator is allowed to reach its optimum given G, and p_q is updated so as to improve the criterion

$$\mathbb{E}_{\boldsymbol{x} \sim p_{data}}[\log D_G^*(\boldsymbol{x})] + \mathbb{E}_{\boldsymbol{x} \sim p_g}[\log(1 - D_G^*(\boldsymbol{x}))]$$

then p_q converges to p_{data}

Proof. Consider $V(G,D)=U(p_g,D)$ as a function of p_g as done in the above criterion. Note that $U(p_g,D)$ is convex in p_g . The subderivatives of a supremum of convex functions include the derivative of the function at the point where the maximum is attained. In other words, if $f(x)=\sup_{\alpha\in\mathcal{A}}f_\alpha(x)$ and $f_\alpha(x)$ is convex in x for every α , then $\partial f_\beta(x)\in\partial f$ if $\beta=\arg\sup_{\alpha\in\mathcal{A}}f_\alpha(x)$. This is equivalent to computing a gradient descent update for p_g at the optimal D given the corresponding G. $\sup_D U(p_g,D)$ is convex in p_g with a unique global optima as proven in Thm 1, therefore with sufficiently small updates of p_g , p_g converges to p_x , concluding the proof.

In practice, adversarial nets represent a limited family of p_g distributions via the function $G(z;\theta_g)$, and we optimize θ_g rather than p_g itself. Using a multilayer perceptron to define G introduces multiple critical points in parameter space. However, the excellent performance of multilayer perceptrons in practice suggests that they are a reasonable model to use despite their lack of theoretical guarantees.

Convergence of Algorithm

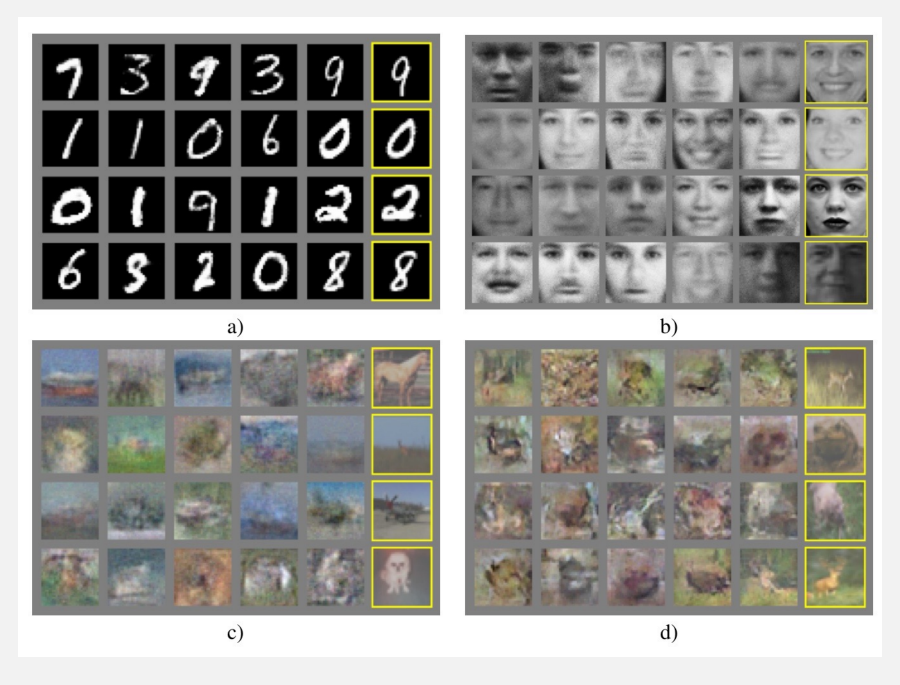
Experiments

Datasets:

MNIST, Toronto Face Database(TFD), CIFAR-10

G: rectifier linear activations(ReLU), sigmoid activations

D: maxout activations, Dropout (Training)



Advantages and Disadvantages

Advantages

- Markov chains are never needed, only backprop is used to obtain gradients
- No inference is needed during learning
- Wide variety of functions can be incorporated into the model
- Model can represent very sharp, even degenerate distributions (than Markov chains)

Disadvantages

- No explicit representation of Pg(x)
- D must synchronized well with G while training

Conclusions and future work

