

# **Accelerated First-Order Optimization Algorithms** for Machine Learning

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ABSTRACT | Numerical optimization serves as one of the pillars of machine learning. To meet the demands of big data applications, lots of efforts have been put on designing theoretically and practically fast algorithms. This article provides a comprehensive survey on accelerated first-order algorithms with a focus on stochastic algorithms. Specifically, this article starts with reviewing the basic accelerated algorithms on deterministic convex optimization, then concentrates on their extensions to stochastic convex optimization, and at last introduces some recent developments on acceleration for nonconvex optimization.

KEYWORDS | Acceleration; convex optimization; deterministic algorithms; machine learning; nonconvex optimization; stochastic algorithms.

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#### I. INTRODUCTION

Many machine learning problems can be formulated as the sum of n loss functions and one regularizer

$$\min_{\mathbf{x} \in \mathbb{R}^p} F(\mathbf{x}) \stackrel{\text{def}}{=} f(\mathbf{x}) + h(\mathbf{x}) \stackrel{\text{def}}{=} \frac{1}{n} \sum_{i=1}^n f_i(\mathbf{x}) + h(\mathbf{x})$$
(1)

where  $f_i(\mathbf{x})$  is the loss function,  $h(\mathbf{x})$  is typically a regularizer, and n is the sample size. Examples of  $f_i(\mathbf{x})$  include  $f_i(\mathbf{x}) = (y_i - \mathbf{A}_i^T \mathbf{x})^2$  for the linear least squared loss and  $f_i(\mathbf{x}) = \log(1 + \exp(-y_i \mathbf{A}_i^T \mathbf{x}))$  for the logistic loss, where  $\mathbf{A}_i \in \mathbb{R}^p$  is the feature vector of the *i*th sample and  $y_i \in \mathbb{R}$ is its target value or label. Representative examples of  $h(\mathbf{x})$ include the  $\ell_2$  regularizer  $h(\mathbf{x}) = (1/2) \|\mathbf{x}\|^2$  and the  $\ell_1$ regularizer  $h(\mathbf{x}) = ||\mathbf{x}||_1$ . Problem (1) covers many famous models in machine learning, for example, support vector machine (SVM) [1], logistic regression [2], LASSO [3], multilayer perceptron [4], and so on.

Optimization plays an indispensable role in machine learning, which involves the numerical computation of the optimal parameters with respect to a given learning model based on the training data. Note that the dimension pcan be very high in many machine learning applications. In such a setting, computing the Hessian matrix of f to use in a second-order algorithm is time-consuming. Thus, first-order optimization methods are usually preferred over high-order ones and they have been the workhorse for a tremendous amount of machine learning applications.

Gradient descent (GD) has been one of the most commonly used first-order method due to its simplicity to implement and low computational cost per iteration. Although practical and effective, GD converges slowly in many applications. To accelerate its convergence, there has

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been a surge of interest in accelerated gradient methods, where "accelerated" means that the convergence rate can be improved without much stronger assumptions or significant additional computational burden. Nesterov [5]–[8] has proposed several accelerated GD (AGD) methods in his celebrated works, which have provable faster convergence rates than the basic GD.

Originating from Nesterov's celebrated works, accelerated first-order methods have become a hot topic in the machine learning community, yielding great success [9]. In machine learning, the sample size n can be extremely large and computing the full gradient in GD or AGD is time-consuming. Therefore, stochastic gradient methods are the coin of the realm to deal with big data, which use only a few randomly chosen samples at each iteration. It motivates the extension of Nesterov's accelerated methods from deterministic optimization to finite-sum stochastic optimization [10]–[20]. Due to the success of deep learning, in recent years, there has been a trend to design and analyze efficient nonconvex optimization algorithms, especially with a focus on accelerated methods [21]–[27].

In this article, we provide a comprehensive survey on the accelerated first-order algorithms. To proceed, we provide some notation and definitions that will frequently be used in this article.

#### A. Notation and Definitions

We use uppercase bold letters to represent matrices, lowercase bold letters for vectors, and nonbold letters for scalars. Let us denote by  $\mathbf{A}_i$  the ith column of  $\mathbf{A}$ ,  $\mathbf{x}_i$  the ith coordinate of  $\mathbf{x}$ , and  $\nabla_i f(\mathbf{x})$  the ith coordinate of  $\nabla f(\mathbf{x})$ . We denote by  $\mathbf{x}^k$  the value of  $\mathbf{x}$  of an algorithm at the kth iteration and  $\mathbf{x}^*$  any optimal solution of problem (1). For scalars, for example,  $\theta$ , we denote by  $\{\theta_k\}_{k=0}^{\infty}$  a sequence of real numbers and by  $\theta_k^2$  the power of  $\theta_k$ .

We study both convex and nonconvex problems in this article.

Definition 1: A function  $f(\mathbf{x})$  is  $\mu$ -strongly convex, meaning that

$$f(\mathbf{y}) \ge f(\mathbf{x}) + \langle \xi, \mathbf{y} - \mathbf{x} \rangle + \frac{\mu}{2} ||\mathbf{y} - \mathbf{x}||^2$$
 (2)

for all  $\mathbf{x}$  and  $\mathbf{y}$ , where  $\xi \in \partial f(\mathbf{x})$  is a subgradient of f. Especially, we allow  $\mu = 0$ , in which case we call  $f(\mathbf{x})$  is nonstrongly convex.

Note that "nonstrongly convex" is frequently used in this article. Therefore, a definition is appropriate. We often assume that the objective function is *L*-smooth, meaning that its gradient cannot change arbitrarily fast.

Definition 2: A function f is L-smooth if it satisfies

$$\|\nabla f(\mathbf{y}) - \nabla f(\mathbf{x})\| < L\|\mathbf{y} - \mathbf{x}\|$$

for all **x** and **y** and some  $L \geq 0$ .

A vital property of an L-smooth function f is

$$f(\mathbf{y}) \le f(\mathbf{x}) + \langle \xi, \mathbf{y} - \mathbf{x} \rangle + \frac{L}{2} ||\mathbf{y} - \mathbf{x}||^2 \quad \forall \mathbf{x}, \mathbf{y}.$$

Classically, the definition of a first-order algorithm in optimization theory is based on an oracle that only returns  $f(\mathbf{x})$  and  $\nabla f(\mathbf{x})$  for a given  $\mathbf{x}$ . Here, we adopt a much more general sense that the oracle also returns the solution of some simple proximal mapping.

*Definition 3:* The proximal mapping of a function h for some given  $\mathbf{z}$  is defined as

$$\operatorname{Prox}_h(\mathbf{z}) = \operatorname*{argmin}_{\mathbf{x} \in \mathbb{R}^p} h(\mathbf{x}) + \frac{1}{2} \|\mathbf{x} - \mathbf{z}\|^2.$$

"Simple" means that the solution can be computed efficiently, and it does not dominate the computation time at each iteration of an algorithm, for example, having a closed solution, which is typical in machine learning. For example, in compressed sensing, we often use  $\text{Prox}_{\lambda \|\cdot\|_1}(\mathbf{z}) = \text{sign}(\mathbf{z}) \max\{0, |\mathbf{z}| - \lambda\}$ . In this article, we consider only algorithms based on the proximal mapping of  $f_i(\mathbf{x})$  or  $h(\mathbf{x})$  in (1), but not that of  $F(\mathbf{x})$ .

In this article, we use iteration complexity to describe the convergence speed of a deterministic algorithm.

Definition 4: For convex problems, we define iteration complexity as the smallest number of iterations needed to find an  $\varepsilon$ -optimal solution within a tolerance  $\varepsilon$  on the error to the optimal objective, that is,  $F(\mathbf{x}) - F(\mathbf{x}^*) \leq \varepsilon$ .

In nonconvex optimization, it is infeasible to describe the convergence speed by  $F(\mathbf{x}) - F(\mathbf{x}^*) \leq \varepsilon$ , since finding the global minima is NP-hard. Alternatively, we use the number of iterations to find an  $\varepsilon$ -approximate stationary point.

*Definition 5:* We say that **x** is an ε-approximate first-order stationary point of problem (1), if it satisfies  $\|\mathbf{x} - \operatorname{Prox}_h(\mathbf{x} - \nabla f(\mathbf{x}))\| \le \varepsilon$ . It reduces to  $\|\nabla f(\mathbf{x})\| \le \varepsilon$  when  $h(\mathbf{x}) = 0$ .

For nonconvex functions, the first-order stationary points can be global minima, local minima, saddle points, or local maxima. Sometimes, it is not enough to find the first-order stationary points and it motivates us to pursue high-order stationary points.

Definition 6: We say that  $\mathbf{x}$  is an  $(\varepsilon, O(\sqrt{\varepsilon}))$ -approximate second-order stationary point of problem (1) with  $h(\mathbf{x})=0$ , if it satisfies  $\|\nabla f(\mathbf{x})\| \leq \varepsilon$  and  $\sigma_{\min}(\nabla^2 f(\mathbf{x})) \geq -O(\sqrt{\varepsilon})$ , where  $\sigma_{\min}(\nabla^2 f(\mathbf{x}))$  means the smallest singular value of the Hessian matrix.

Intuitively speaking,  $\|\nabla f(\mathbf{x})\| = 0$  and  $\nabla^2 f(\mathbf{x}) \succeq 0$  means that  $\mathbf{x}$  is either a local minima or a higher-order saddle point. Since higher-order saddle points do not exist for many machine learning problems and all local minima are global minima, for example, in matrix sensing [28], matrix completion [29], robust PCA [30], and deep neural networks [31], [32], it is enough to find second-order stationary points for these problems.

For stochastic algorithms, to emphasize the dependence on the sample size n, we use gradient complexity to describe the convergence speed.

Definition 7: The gradient complexity of a stochastic algorithm is defined as the number of accessing the individual gradients for searching an  $\varepsilon$ -optimal solution or an  $\varepsilon$ -approximate stationary point in expectation, that is, replacing  $F(\mathbf{x})$ ,  $\|\nabla f(\mathbf{x})\|$ , and  $\sigma_{\min}(\nabla^2 f(\mathbf{x}))$  by  $\mathbb{E}[F(\mathbf{x})]$ ,  $\mathbb{E}[\|\nabla f(\mathbf{x})\|]$ , and  $\mathbb{E}[\sigma_{\min}(\nabla^2 f(\mathbf{x}))]$  in the above definitions,

Finally, we define the Bregman distance. The most commonly used Bregman distance is  $D(\mathbf{x}, \mathbf{y}) = (1/2) \|\mathbf{x} - \mathbf{y}\|^2$ .

Definition 8: Bregman distance is defined as

$$D_{\varphi}(\mathbf{u}, \mathbf{v}) = \varphi(\mathbf{u}) - (\varphi(\mathbf{v}) + \langle \hat{\nabla} \varphi(\mathbf{v}), \mathbf{u} - \mathbf{v} \rangle)$$
(3)

for strongly convex  $\varphi$  and  $\hat{\nabla}\varphi(\mathbf{v}) \in \partial\varphi(\mathbf{v})$ .

# II. BASIC ACCELERATED DETERMINISTIC ALGORITHMS

In this section, we discuss the speedup guarantees of the basic accelerated gradient methods over the basic GD for deterministic convex optimization.

#### A. Gradient Descent

GD and its proximal variant have been one of the most commonly used first-order deterministic method. The latter one consists of the following iterations:

$$\mathbf{x}^{k+1} = \text{Prox}_{\eta h}(\mathbf{x}^k - \eta \nabla f(\mathbf{x}^k))$$

where we assume that f is L-smooth.  $\eta$  is the step size and it is usually set to 1/L. When the objective  $f(\mathbf{x})$  in (1) is L-smooth and  $\mu$ -strongly convex, and  $h(\mathbf{x})$  is convex, GD and its proximal variant converge linearly [33], which is described as

$$F(\mathbf{x}^k) - F(\mathbf{x}^*) \leq \left(1 - \frac{\mu}{L}\right)^k L \|\mathbf{x}^0 - \mathbf{x}^*\|^2 = O\left(\left(1 - \frac{\mu}{L}\right)^k\right).$$

In other words, the iteration complexity of GD is  $O((L/\mu)\log(1/\varepsilon))$  to find an  $\varepsilon$ -optimal solution.

When  $f(\mathbf{x})$  is smooth and nonstrongly convex, GD obtains only a sublinear rate [33] of

$$F(\mathbf{x}^k) - F(\mathbf{x}^*) \le \frac{L}{2(k+1)} ||\mathbf{x}^0 - \mathbf{x}^*||^2 = O(1/k).$$

In this case, the iteration complexity of GD becomes  $O(L/\varepsilon)$ .

# B. Heavy-Ball Method

The convergence speed of GD for strongly convex problems is determined by the constant  $L/\mu$ , which is

known as the condition number of  $f(\mathbf{x})$ , and it is always greater or equal to 1. When the condition number is very large, that is, the problem is ill-conditioned, GD converges slowly. Accelerated methods can speed up over GD significantly for ill-conditioned problems.

Polyak's heavy-ball method [34] was the first accelerated gradient method. It counts for the history of iterates when computing the next iterate. The next iterate depends not only on the current iterate, but also on the previous ones. The proximal variant of the heavy-ball method [35] is

$$\mathbf{x}^{k+1} = \operatorname{Prox}_{\eta h}(\mathbf{x}^k - \eta \nabla f(\mathbf{x}^k) + \beta(\mathbf{x}^k - \mathbf{x}^{k-1}))$$
 (4)

where  $\eta = 4/(\sqrt{L} + \sqrt{\mu})^2$  $(\sqrt{L} - \sqrt{\mu})^2/(\sqrt{L} + \sqrt{\mu})^2$ . When  $f(\mathbf{x})$  is L-smooth and  $\mu$ -strongly convex and  $h(\mathbf{x})$  is convex, and moreover,  $f(\mathbf{x})$  is twice continuously differentiable, the heavy-ball method and its proximal variant have the following local accelerated convergence rate [35]:

$$F(\mathbf{x}^k) - F(\mathbf{x}^*) \leq O\left(\left(\frac{\sqrt{L} - \sqrt{\mu}}{\sqrt{L} + \sqrt{\mu}}\right)^k\right) \leq O\left(\left(1 - \sqrt{\frac{\mu}{L}}\right)^k\right).$$

So, the iteration complexity of the heavy-ball method is  $O(\sqrt{L/\mu}\log(1/\varepsilon))$ , which is significantly lower than that of the basic GD when  $L/\mu$  is large. The twice continuous differentiability is necessary to ensure the convergence. Otherwise, the heavy-ball method may fail to converge even for strongly convex problems [36].

When the strong convexity assumption is absent, currently only the  $O(L/\varepsilon)$  iteration complexity is proved for the heavy-ball method [37], which is the same as the basic GD. Theoretically, it is unclear whether the O(1/k) rate is tight. Ghadimi et al. [37] numerically observed that O(1/k) is an accurate convergence rate estimate for the heavy-ball method. Next, we introduce Nesterov's basic accelerated gradient methods to further speedup the convergence for nonstrongly convex problems.

### C. Nesterov's Accelerated Gradient Method

Nesterov's accelerated gradient methods have faster convergence rates than the basic GD for both strongly convex and nonstrongly convex problems. In its simplest form, the proximal variant of Nesterov's AGD [38] takes the form

$$\mathbf{y}^k = \mathbf{x}^k + \beta_k (\mathbf{x}^k - \mathbf{x}^{k-1}) \tag{5a}$$

$$\mathbf{x}^{k+1} = \operatorname{Prox}_{nh}(\mathbf{y}^k - \eta \nabla f(\mathbf{y}^k)). \tag{5b}$$

Physically, AGD first adds a momentum, that is,  $\mathbf{x}^k - \mathbf{x}^{k-1}$ , to the current point  $\mathbf{x}^k$  to generate an extrapolated point  $\mathbf{y}^k$  and then performs a proximal GD step at  $\mathbf{y}^k$ . Similar to the heavy-ball method, the iteration complexity of (5a) and (5b) is  $O(\sqrt{L/\mu}\log(1/\varepsilon))$  for problem (1) with L-smooth and  $\mu$ -strongly convex  $f(\mathbf{x})$  and convex  $h(\mathbf{x})$ , by setting  $\eta = 1/L$ ,  $\beta_k \equiv (\sqrt{L} - \sqrt{\mu})/(\sqrt{L} + \sqrt{\mu})$ , and  $\mathbf{x}^0 = \mathbf{x}^{-1}$  [33]. However, it does not need the assumption of twice continuous differentiability of  $f(\mathbf{x})$ .

Better than the heavy-ball method, for smooth and nonstrongly convex problems, AGD has a faster sublinear rate described as

$$F(\mathbf{x}^k) - F(\mathbf{x}^*) \le O(1/k^2)$$

and the iteration complexity is improved to  $O(\sqrt{L/\varepsilon})$ . One often sets  $\beta_k = \theta_k (1-\theta_{k-1})/\theta_{k-1}$  for nonstrongly convex problems, where the positive sequence  $\{\theta_k\}_{k=0}^\infty$  is obtained by solving equation  $\theta_k^2 = (1-\theta_k)\theta_{k-1}^2$ , which is initialized by  $\theta_0 = 1$ . Sometimes, one sets  $\beta_k = (k-1)/(k+2)$  for simplicity.

Physically, the acceleration can be interpreted as adding momentum to the iterates. Also, Su *et al.* [39] derived a second-order ordinary differential equation to model scheme (5a) and (5b), Safavi *et al.* [40] analyzed it via the notion of integral quadratic constraints [36] from the robust control theory and Wibisono *et al.* [41] further explained the mechanism of acceleration from a continuous-time variational point of view.

#### D. Other Variants and Extensions

Besides the basic AGD (5a) and (5b), Nesterov [6]–[8] and Nesterov *et al.* [42] also proposed several other accelerated methods, and Tseng [43] further provided a unified analysis. We briefly introduce the method in [6], which is easier to extend to many other variants than (5a) and (5b). These variants include accelerated variance reduction (VR) [10], accelerated randomized coordinate descent [15], [16], and accelerated asynchronous algorithm [44]. This method consists of the following iterations:

$$\mathbf{y}^k = (1 - \theta_k)\mathbf{x}^k + \theta_k \mathbf{z}^k \tag{6a}$$

$$\mathbf{z}^{k+1} = \operatorname{Prox}_{h/(L\theta_k)} \left( \mathbf{z}^k - \frac{1}{L\theta_k} \nabla f(\mathbf{y}^k) \right)$$
 (6b)

$$\mathbf{x}^{k+1} = (1 - \theta_k)\mathbf{x}^k + \theta_k\mathbf{z}^{k+1}$$
 (6c)

where  $\theta_k$  is the same as that in (5a) and (5b), and we initialize  $\mathbf{z}^0 = \mathbf{x}^0$ . Note that (5a) and (5b) and (6a)–(6c) produce the same iterates  $\mathbf{y}^k$  and  $\mathbf{x}^k$  when  $h(\mathbf{x}) = 0$ . To explain the mechanism of acceleration, Allen-Zhu and Orecchia [45] explicated (6a)–(6c) by linear coupling [step (6a)] of GD [step (6c)] and mirror descent [step (6b)]. Lan and Zhou [20] viewed (6a)–(6c) as an iterative buyer–supplier game by rewriting it in an equivalent primal–dual form [46], [47].

Motivated by Nesterov's celebrated work, some researchers have proposed other accelerated methods. Bubeck *et al.* [48] proposed a geometric descent method, which has a simple geometric interpretation of accele-

ration. Drusvyatskiy et al. [49] explained the geometric descent from the perspective of optimal average of quadratic lower models, which is related to Nesterov's estimate sequence technique [33]. However, the methods in [48] and [49] need a line-search step. They minimize  $f(\mathbf{x})$  exactly on the line between two points  $\mathbf{x}$  and  $\mathbf{y}$ . Thus, their methods are not rigorously "first-order" methods. Drori and Teboulle [50] proposed a numerical procedure for computing optimal tuning coefficients in a class of first-order algorithms, including Nesterov's AGD. Motivated by Drori and Teboulle [50], Kim and Fessler [51] introduced several new optimized first-order methods whose coefficients are analytically found. Lan [52], Lan and Zhou [53], and Lan and Ouyang [54] extended the accelerated methods to some complex composite convex optimization and structured convex optimization via the gradient sliding technique, where an inner loop is used to skip some computations from time to time. The acceleration technique has also been used to solve linearly constrained problems [55]-[59]. However, many methods for constrained problems [46], [47], [60]-[64] need to solve an optimization subproblem exactly, thus they are not "first-order" methods either.

#### E. Lower Bound

Can we find algorithms faster than AGD? Better yet, how fast can we solve problem (1), or its simplified case

$$\min_{\mathbf{x} \in \mathbb{R}^p} f(\mathbf{x}) \tag{7}$$

to some accuracy  $\varepsilon$ , using methods only based on the information of  $\nabla f(\mathbf{x})$ ? A few existing bounds can answer these questions. The first lower bounds for first-order optimization algorithms were given in [65] and then were extended in [33]. We introduce the widely used conclusion in [33]. Consider any iterative first-order method generating a sequence of points  $\{\mathbf{x}^t\}_{t=0}^k$  such that

$$\mathbf{x}^k \in \mathbf{x}^0 + \operatorname{Span}\{\nabla f(\mathbf{x}^0), \dots, \nabla f(\mathbf{x}^{k-1})\}.$$
 (8)

Nesterov [33] constructed a special L-smooth and  $\mu$ -strongly convex function  $f(\mathbf{x})$  such that for any sequence satisfying (8), we have

$$f(\mathbf{x}^k) - f(\mathbf{x}^*) \ge \frac{\mu}{2} \left( \frac{\sqrt{L} - \sqrt{\mu}}{\sqrt{L} + \sqrt{\mu}} \right)^{2k} \|\mathbf{x}^0 - \mathbf{x}^*\|^2.$$

It means that any first-order method satisfying (8) needs at least  $O(\sqrt{L/\mu}\log(1/\varepsilon))$  iterations to achieve an  $\varepsilon$ -optimal solution for the class of L-smooth and  $\mu$ -strongly convex problems. Recalling the upper bound given in Section II-C, we can see that it matches this lower bound. Thus, Nesterov's AGDs are optimal and they cannot be further accelerated up to constants. When the strong convexity

Table 1 Iteration Complexity Comparisons Between GD, the Heavy-Ball Method, and AGD, as Well as the Lower Bounds

Method	Strongly convex	Non-strongly convex	
GD	$O\left(\frac{L}{\mu}\log\frac{1}{\varepsilon}\right)$ [33]	$O\left(\frac{L}{\varepsilon}\right)$ [33]	
heavy-ball	$O\left(\sqrt{\frac{L}{\mu}}\log\frac{1}{\varepsilon}\right)$ [35]	$O\left(\frac{L}{\varepsilon}\right)$ [37]	
AGD	$O\left(\sqrt{\frac{L}{\mu}}\log\frac{1}{\varepsilon}\right)$ [8], [33]	$O\left(\sqrt{\frac{L}{\varepsilon}}\right)$ [8], [33], [38], [51]	
Lower Bounds	$O\left(\sqrt{\frac{L}{\mu}}\log\frac{1}{\varepsilon}\right)$ [33], [66], [67]	$O\left(\sqrt{\frac{L}{\varepsilon}}\right)$ [33], [66], [67]	

is absent, Nesterov [33] constructed another L-smooth convex function  $f(\mathbf{x})$  such that for any method satisfying (8), we have

$$f(\mathbf{x}^k) - f(\mathbf{x}^*) \ge \frac{3L}{32(k+1)^2} \|\mathbf{x}^0 - \mathbf{x}^*\|^2$$
$$\|\mathbf{x}^k - \mathbf{x}^*\|^2 \ge \frac{1}{32} \|\mathbf{x}^0 - \mathbf{x}^*\|^2.$$

The iteration number k for the counterexample in [33] depends on the dimension p of the problem, for example, k should satisfy  $k \leq (1/2)(p-1)$  for nonstrongly convex problems. Arjevani and Shamir [66] and Arjevani et al. [68] proposed a different framework to establish the same lower bounds as [33], but, the iteration number in [66] and [68] is dimension-independent. When considering the composite problem (1), we can use the results in [67] to give the same lower bounds as [33] for first-order methods that are only based on the information of  $\nabla f(\mathbf{x})$  and  $Prox_h(\mathbf{z})$ . Although Woodworth and Srebro [67] studied the finite-sum problem, their conclusion can be used to (1) as long as  $f(\mathbf{x}) \neq 0$ ,  $h(\mathbf{x}) \neq 0$ , and  $f(\mathbf{x}) \neq h(\mathbf{x})$ .

For better comparison of different methods, we list the iteration complexities as well as the lower bounds in Table 1.

# III. ACCELERATED STOCHASTIC ALGORITHMS

In machine learning, people often encounter big data with extremely large n in problem (1). Computing the full gradient of  $f(\mathbf{x})$  in GD and AGD might be expensive. Stochastic gradient algorithms might be the most common way to cope with big data. They sample, in each iteration, one or several gradients from individual functions as an estimator of the full gradient of f. For example, consider the standard proximal stochastic GD (SGD), which uses one stochastic gradient at each iteration and proceeds as follows:

$$\mathbf{x}^{k+1} = \text{Prox}_{\eta_k h}(\mathbf{x}^k - \eta_k \nabla f_{i_k}(\mathbf{x}^k))$$

where  $\eta_k$  denotes the step size and  $i_k$  is an index randomly sampled from  $\{1, \ldots, n\}$  at iteration k. SGD often suffers from slow convergence. For example, when the objective is L-smooth and  $\mu$ -strongly convex, SGD obtains only a sublinear rate [69] of

$$\mathbb{E}[F(\mathbf{x}^k)] - F(\mathbf{x}^*) \le O(1/k).$$

In contrast, GD has the linear convergence. In Sections III-A-III-C, we introduce several techniques to accelerate SGD. In particular, we discuss how Nesterov's acceleration works in stochastic optimization with finite n in problem (1), which is often called the finite-sum problem.

#### A. VR and Its Acceleration

The main challenge for SGD is the noise of the randomly drawn gradients. The variance of the noisy gradient will never go to zero even if  $\mathbf{x}^k \rightarrow \mathbf{x}^*$ . As a result, one has to gradually cut down the step size in SGD to guarantee convergence, which brings down the convergence. A technique called VR [70] was designed to reduce the negative effect of noise. For finite-sum objective functions, the VR technique reduces the variance to zero through the updates. The first VR method might be stochastic average gradient (SAG) [71], which uses the sum of the latest individual gradients as an estimator of the descent direction. It requires O(np) memory storage and uses a biased gradient estimator. Stochastic variance reduced gradient (SVRG) [70] reduces the memory cost to O(p) and uses an unbiased gradient estimator. Later, SAGA [72] improves SAG by using an unbiased update via the technique of SVRG. Other VR methods can be found in [73]–[78].

We take SVRG [79] as an example, which is relatively simple and easy to implement. SVRG maintains a snapshot vector  $\tilde{\mathbf{x}}^s$  after every m SGD iterations and keeps the gradient of the averages  $\mathbf{g}^s = \nabla f(\tilde{\mathbf{x}}^s)$ . Then, it uses  $\tilde{\nabla} f(\mathbf{x}^k) = \nabla f_{i_k}(\mathbf{x}^k) - \nabla f_{i_k}(\tilde{\mathbf{x}}^s) + \mathbf{g}^s$  as the descent direction at every SGD iterations and the expectation  $\mathbb{E}_{i_k}[\tilde{\nabla}f(\mathbf{x}^k)] =$  $\nabla f(\mathbf{x}^k)$ . Moreover, the variance of the estimated gradient  $\tilde{\nabla} f(\mathbf{x}^{s,k})$  now can be upper bounded by the distance from the snapshot vector to the latest variable, that is,  $\mathbb{E}[\|\tilde{\nabla}f(\mathbf{x}^k) - \nabla f(\mathbf{x}^k)\|^2] \leq L\|\mathbf{x}^k - \tilde{\mathbf{x}}^s\|^2$ , which is a crucial property of SVRG to guarantee the reduction of variance. Algorithm 1 gives the details of SVRG.

# Algorithm 1 SVRG

Input 
$$\tilde{\mathbf{x}}^0$$
,  $m = O(L/\mu)$ , and  $\eta = O(1/L)$ .

for  $s = 0, 1, 2, \cdots$  do

 $\mathbf{x}^0 = \tilde{\mathbf{x}}^s$ ,

 $\mathbf{g}^s = \nabla f(\tilde{\mathbf{x}}^s)$ ,

for  $k = 0, \cdots, m$  do

Randomly sample  $i_k$  from  $\{1, \dots, n\}$ ,

 $\tilde{\nabla} f(\mathbf{x}^k) = \nabla f_{i_k}(\mathbf{x}^k) - \nabla f_{i_k}(\tilde{\mathbf{x}}^s) + \mathbf{g}^s$ ,

 $\mathbf{x}^{k+1} = \operatorname{Prox}_{\eta h} \left(\mathbf{x}^k - \eta \tilde{\nabla} f(\mathbf{x}^k)\right)$ ,

end for

 $\tilde{\mathbf{x}}^{s+1} = \frac{1}{m} \sum_{k=1}^m \mathbf{x}^k$ ,

end for

For  $\mu$ -strongly convex problem (1) with L-smooth  $f_i(\mathbf{x})$ , SVRG needs  $O((L/\mu)\log(1/\varepsilon))$  inner iterations to reach an  $\varepsilon$ -optimal solution in expectation. Each inner iteration needs to evaluate two stochastic gradients, while each outer iteration needs additional n individual gradient evaluations to compute  $\mathbf{g}^s$ . Thus, the gradient complexity of SVRG is  $O((n+L/\mu)\log(1/\varepsilon))$ . Recall that GD has the gradient complexity of  $O((nL/\mu)\log(1/\varepsilon))$ , since it needs n individual gradient evaluations at each iteration. Thus, SVRG is superior to GD when  $L/\mu > 1$ .

With the VR technique in hand, one can fuse it with Nesterov's acceleration technique to further accelerate stochastic algorithms (see [10]–[13], [80], and [81]). We take Katyusha [10] as an example. Katyusha builds upon the combination of (6a)–(6c) and SVRG. Different from (6a) and (6c), Katyusha further introduces a "negative momentum" with additional  $\tau'\bar{\mathbf{x}}^s$  in (9a) and (9d), which prevents the extrapolation term from being far from the snapshot vector. Algorithm 2 gives the details of Katyusha.

# Algorithm 2 Katyusha

Input 
$$\mathbf{x}^0 = \mathbf{z}^0 = \tilde{\mathbf{x}}^0$$
,  $m = n$ ,  $\tau = \min\{\sqrt{\frac{n\mu}{3L}}, \frac{1}{2}\}$ ,  $\tau' = \frac{1}{2}$ ,  $\eta = O(\frac{1}{L})$ , and  $\tau'' = \frac{\mu}{3\tau L} + 1$ .

for  $s = 0, 1, 2, \cdots$  do

 $\mathbf{g}^s = \nabla f(\tilde{\mathbf{x}}^s)$ 

for  $k = 0, \cdots, m$  do

 $\mathbf{y}^k = \tau \mathbf{z}^k + \tau' \tilde{\mathbf{x}}^s + (1 - \tau - \tau') \mathbf{x}^k$ , (9a)

Randomly sample  $i_k$  from  $\{1, \dots, n\}$ ,

 $\tilde{\nabla} f(\mathbf{y}^k) = \nabla f_{i_k}(\mathbf{y}^k) - \nabla f_{i_k}(\tilde{\mathbf{x}}^s) + \mathbf{g}^s$ , (9b)

 $\mathbf{z}^{k+1} = \operatorname{Prox}_{\eta h/\tau} \left(\mathbf{z}^k - \eta/\tau \tilde{\nabla} f(\mathbf{y}^k)\right)$ , (9c)

 $\mathbf{x}^{k+1} = \tau \mathbf{z}^{k+1} + \tau' \tilde{\mathbf{x}}^s + (1 - \tau - \tau') \mathbf{x}^k$ , (9d)

$$\begin{array}{l} \text{end for} \\ \tilde{\mathbf{x}}^{s+1} = \left(\sum_{k=0}^{m-1} (\tau'')^k\right)^{-1} \sum_{k=0}^{m-1} (\tau'')^k \mathbf{x}^k, \\ \mathbf{z}^0 = \mathbf{x}^m, \, \mathbf{x}^0 = \mathbf{x}^m, \\ \text{end for} \end{array}$$

For problems with smooth and convex  $f_i(\mathbf{x})$  and  $\mu$ -strongly convex  $h(\mathbf{x})$ , the gradient complexity of Katyusha is  $O((n+\sqrt{nL/\mu})\log(1/\varepsilon))$ . When  $n \leq O(L/\mu)$ , Katyusha further accelerates SVRG. Comparing SVRG with Katyusha, we can see that the only difference is that Katyusha uses the mechanism of AGD in (6a)–(6c). Thus, Nesterov's acceleration technique also takes effect in finite-sum stochastic optimization.

We now describe several extensions of Katyusha with some advanced topics.

1) Loopless Katyusha: Both SVRG and Katyusha have double loops, which make them a little complex to analyze and implement. To remedy the double loops, Kovalev *et al.* [12] proposed a loopless SVRG and Katyusha for the

simplified problem

$$\min_{\mathbf{x} \in \mathbb{R}^p} f(\mathbf{x}) \stackrel{\text{def}}{=} \frac{1}{n} \sum_{i=1}^n f_i(\mathbf{x})$$
 (10)

with smooth and convex  $f_i(\mathbf{x})$  and strongly convex  $f(\mathbf{x})$ . Specifically, at each iteration, with a small probability 1/n, the methods update the snapshot vector and perform a full pass over data to compute the average gradient. With probability 1-1/n, the methods use the previous snapshot vector. The loopless SVRG and Katyusha enjoy the same gradient complexities as the original methods. We take the loopless SVRG as an example, which consists of the following steps at each iteration:

$$\tilde{\nabla}f(\mathbf{x}^k) = \nabla f_{i_k}(\mathbf{x}^k) - \nabla f_{i_k}(\tilde{\mathbf{x}}^k) + \nabla f(\tilde{\mathbf{x}}^k)$$
(11a)

$$\mathbf{x}^{k+1} = \mathbf{x}^k - \eta \tilde{\nabla} f(\mathbf{x}^k) \tag{11b}$$

$$\tilde{\mathbf{x}}^{k+1} = \begin{cases} \mathbf{x}^k, & \text{with probability } 1/n, \\ \tilde{\mathbf{x}}^k, & \text{with probability } 1 - 1/n. \end{cases}$$
 (11c)

When replacing (11a) and (11b) by the following steps, we obtain the loopless Katyusha:

$$\begin{split} &\mathbf{y}^k = \tau \mathbf{z}^k + \tau' \tilde{\mathbf{x}}^k + (1 - \tau - \tau') \mathbf{x}^k \\ \tilde{\nabla} f(\mathbf{y}^k) &= \nabla f_{i_k}(\mathbf{y}^k) - \nabla f_{i_k}(\tilde{\mathbf{x}}^k) + \nabla f(\tilde{\mathbf{x}}^k) \\ &\mathbf{z}^{k+1} = \frac{1}{\alpha \mu/L + 1} \left( \frac{\alpha \mu}{L} \mathbf{y}^k + \mathbf{z}^k - \frac{\alpha}{L} \tilde{\nabla} f(\mathbf{y}^k) \right) \\ &\mathbf{x}^{k+1} = \tau \mathbf{z}^{k+1} + \tau' \tilde{\mathbf{x}}^s + (1 - \tau - \tau') \mathbf{x}^k \end{split}$$

where we set  $\tau = \min\{(2n\mu/(3L))^{1/2}, 1/2\}, \tau' = 1/2$ , and  $\alpha = 2/(3\tau)$ .

- 2) Nonstrongly Convex Problems: When the strong convexity assumption is absent, the gradient complexities of SGD and SVRG are  $O(1/\varepsilon^2)$  [82] and  $O(n \log(1/\varepsilon) + L/\varepsilon)$  [83], respectively. Katyusha improves the complexity of SVRG to  $O(n((F(\mathbf{x}^0)) F(\mathbf{x}^*)/\varepsilon)^{1/2} + (nL\|\mathbf{x}^0 - \mathbf{x}^*\|^2/\varepsilon)^{1/2})$  [10], [11]. This gradient complexity is not more advantageous over Nesterov's full batch AGD since they all need  $O(n/\sqrt{\varepsilon})$ individual gradient evaluations. When applying reductions to extend the algorithms designed for smooth and strongly convex problems to nonstrong convex ones, for example, the HOOD framework [84], the gradient complexity of Katyusha can be further improved to  $O(n\log(1/\varepsilon) + \sqrt{nL/\varepsilon})$ , which is  $\sqrt{n}$  times faster than the full batch AGD when high precision is required. On the other hand, Lan et al. [13] proposed a unified VR accelerated gradient method, which employs a direct acceleration scheme instead of employing any reduction to obtain the desired gradient complexity of  $O(n \log n + \sqrt{nL/\varepsilon}).$
- 3) Universal Catalyst Acceleration for First-Order Convex Optimization: Another way to accelerate SVRG is to use

the universal catalyst [85], which is a unified framework to accelerate first-order methods. It builds upon the accelerated proximal point method with inexactly computed proximal mapping. Analogous to (5a) and (5b), the catalyst takes the following outer iterations:

$$\mathbf{y}^k = \mathbf{x}^k + \beta_k (\mathbf{x}^k - \mathbf{x}^{k-1}) \tag{12a}$$

$$\mathbf{x}^k \approx \underset{\mathbf{x} \in \mathbb{R}^p}{\operatorname{argmin}} \{ G^k(\mathbf{x}) \stackrel{\text{def}}{=} F(\mathbf{x}) + \frac{\gamma}{2} \|\mathbf{x} - \mathbf{y}^k\|^2 \}$$
 (12b)

where  $\beta_k = (\sqrt{\gamma + \mu} - \sqrt{\mu})/(\sqrt{\gamma + \mu} + \sqrt{\mu})$  for strongly convex problems, and it updates in the same way as that in (5a) and (5b) for nonstrongly convex ones. We can use any linearly convergent method that is only based on the information of  $\nabla f_i(\mathbf{x})$  and  $\text{Prox}_h(\mathbf{z})$  to approximately solve the subproblem in (12b). The subproblem often has a good condition number and so can be solved efficiently to a high precision. Take SVRG as an example. When we use SVRG to solve the subproblem, the catalyst accelerates SVRG to the gradient complexity of O((n + $\sqrt{nL/\mu}$ ) log $(L/\mu)$  log $(1/\varepsilon)$ ) for strongly convex problems and  $O(\sqrt{nL/\varepsilon \log(1/\varepsilon)})$  for nonstrongly convex ones, by setting  $\gamma = (L - \mu)/(n+1) - \mu$  and the inner iteration number in step (12b) as  $O((n + (L + \gamma)/(\mu + \gamma)) \log(1/\varepsilon))$ with  $\mu \geq 0$  and  $L \geq (n+2)\mu$ . Besides SVRG, the catalyst can also accelerate other methods, for example, SAG and SAGA. The price for generality is that the gradient complexities of the catalyst have an additional poly-logarithmic factor compared with those of Katyusha.

- 4) Individually Nonconvex: Some problems in machine learning can be written as minimizing strongly convex functions that are finite average of nonconvex ones [75], [77]. That is, each  $f_i(\mathbf{x})$  in problem (10) is L-smooth and may be nonconvex, but their average  $f(\mathbf{x})$  is  $\mu$ -strongly convex. Examples include the core machinery for PCA and SVD. SVRG can also be used to solve this problem with the gradient complexity of  $O((n + \sqrt{nL/\mu}) \log(1/\varepsilon))$ [80]. Allen-Zhu [80] further proposed a method named KatyushaX to improve the gradient complexity to O((n + $n^{3/4}\sqrt{L/\mu}\log(1/\varepsilon)$ .
- 5) Lower Complexity Bounds: Similar to the lower bounds for the class of deterministic first-order algorithms, there are also lower bounds for the randomized first-order methods for finite-sum problems. Considering problem (10), Woodworth and Srebro [67] proved that for any firstorder algorithm that is only based on the information of  $\nabla f_i(\mathbf{x})$  and  $\operatorname{Prox}_{f_i}(\mathbf{z})$ , the lower bound for L-smooth and  $\mu$ -strongly convex problems is  $O((n + \sqrt{nL/\mu})\log(1/\varepsilon))$ . When the strong convexity is absent, the lower bound becomes  $O(n+\sqrt{nL/\varepsilon})$ . For better comparison, we list the upper and lower bounds in Table 2.
- 6) Application to Distributed Optimization: VR has also been applied to distributed optimization. Classical distributed algorithms include the distributed GD (DGD) [86], EXTRA [87], the gradient-tracking-based methods

Table 2 Gradient Complexity Comparisons Between SGD, SVRG, and Accelerated VR Methods (AccVR), as Well as the Lower Bounds

Method	Smooth and Strongly Convex	Smooth and Non-strongly Convex
SGD	$O\left(\frac{1}{\varepsilon}\right)$ [69]	$O\left(\frac{1}{\varepsilon^2}\right)$ [82]
SVRG	$O\left(\left(n + \frac{L}{\mu}\right) \log \frac{1}{\varepsilon}\right)$	$O\left(n\log\frac{1}{\varepsilon} + \frac{L}{\varepsilon}\right)$
	[12], [70]–[72]	[83]
AccVR	$O\left(\left(n + \sqrt{\frac{nL}{\mu}}\right)\log\frac{1}{\varepsilon}\right)$	$O\left(n\log n + \sqrt{\frac{nL}{\varepsilon}}\right)$
	[10]–[13]	[13]
Lower bounds	$O\left(\left(n + \sqrt{\frac{nL}{\mu}}\right)\log\frac{1}{\varepsilon}\right)$ [67]	$O\left(n + \sqrt{\frac{nL}{\varepsilon}}\right)$ [67]

[88]-[91], and distributed SGD (DSGD) [92], [93]. To further improve the convergence of stochastic distributed algorithms, Mokhtari and Ribeiro [94] combined EXTRA with SAGA, Xin et al. [95] combined gradient tracking with SAGA, and Xin et al. [95] and Li et al. [96] implemented gradient tracking in SVRG (see [97] for a detailed review). It is an interesting work to implement accelerated VR in distributed optimization in the future.

# B. Stochastic Coordinate Descent (SCD) and Its Acceleration

In problem (1), we often assume that  $f_i(\mathbf{x})$  is smooth and allow  $h(\mathbf{x})$  to be nondifferentiable. However, not all machine learning problems satisfy this assumption. The typical example is SVM, which can be formulated as

$$\min_{\mathbf{x} \in \mathbb{R}^p} F(\mathbf{x}) \stackrel{\text{def}}{=} \frac{1}{n} \sum_{i=1}^n \underbrace{\max\{0, 1 - y_i \mathbf{A}_i^T \mathbf{x}\}}_{f_i(\mathbf{x})} + \underbrace{\frac{\mu}{2} \|\mathbf{x}\|^2}_{h(\mathbf{x})}. \tag{13}$$

We can see that each  $f_i(\mathbf{x})$  is convex but nondifferentiable, and  $h(\mathbf{x})$  is smooth and strongly convex. In practice, we often minimize the negative of the dual of (13), written

$$\min_{\mathbf{u} \in \mathbb{R}^n} D(\mathbf{u}) \stackrel{\text{def}}{=} \frac{1}{2\mu} \left\| \frac{\widetilde{\mathbf{A}}\mathbf{u}}{n} \right\|^2 - \frac{1}{n} \sum_{i=1}^n \mathbf{u}_i + \frac{1}{n} \sum_{i=1}^n I_{[0,1]}(\mathbf{u}_i)$$
 (14)

where  $\widetilde{\mathbf{A}}_i = y_i \mathbf{A}_i$ , and  $I_{[0,1]}(u) = 0$  if  $0 \le u \le 1$ , and  $\infty$ , otherwise. Motivated by (14), we consider the following problem in this section:

$$\min_{\mathbf{u} \in \mathbb{R}^n} D(\mathbf{u}) \stackrel{\text{def}}{=} \Phi(\mathbf{u}) + \sum_{i=1}^n \Psi_i(\mathbf{u}_i). \tag{15}$$

We assume that  $\Phi(\mathbf{u})$  satisfies the coordinate-wise smooth condition  $\|\nabla_i \Phi(\mathbf{u}) - \nabla_i \Phi(\mathbf{v})\| \le L_i \|\mathbf{u} - \mathbf{v}\|$  for any  $\mathbf{u}$  and **v** satisfying  $\mathbf{u}_i = \mathbf{v}_i, \forall j \neq i$ . We also assume that  $\Phi(\mathbf{u})$ is  $\mu$ -strongly convex with respect to norm  $\|\cdot\|_L$ , that is, replacing  $\|\mathbf{y} - \mathbf{x}\|^2$  in (2) by  $\|\mathbf{y} - \mathbf{x}\|_L^2 = \sum_{i=1}^n L_i(\mathbf{y}_i - \mathbf{y}_i)$  $\mathbf{x}_i)^2$ . We require  $\Psi_i(u)$  to be convex but can be nondifferentiable. Take problem (14) as an example,  $L_i =$  $\|\mathbf{A}_i\|^2/(n^2\mu)$ , but the first term in (14) is not strongly convex when n > p.

SCD is a popular method to solve problem (15). It first computes the partial derivative with respect to one randomly chosen variable and then updates this variable by a coordinate-wise GD while keeping the other variables unchanged. SCD is sketched as follows:

$$\mathbf{u}_{i_k}^{k+1} = \operatorname*{argmin}_{u} \left( \Psi_{i_k}(u) + \langle \nabla_{i_k} \Phi(\mathbf{u}^k), u \rangle + \frac{L_{i_k}}{2} |u - \mathbf{u}_{i_k}^k|^2 \right)$$

where  $i_k$  is randomly sampled form  $\{1,\ldots,n\}$ . In SCD, we often assume that the proximal mapping of  $\Psi_i(u)$  can be efficiently computed with a closed solution. Also, we need to compute  $\nabla_{i_k}\Phi(\mathbf{u}^k)$  efficiently. Take problem (14), for example, by keeping track of  $\mathbf{s}^k = \widetilde{\mathbf{A}}\mathbf{u}^k$  and updating  $\mathbf{s}^{k+1}$  by  $\mathbf{s}^k - \widetilde{\mathbf{A}}_{i_k}\mathbf{u}^k_{i_k} + \widetilde{\mathbf{A}}_{i_k}\mathbf{u}^{k+1}_{i_k}$ , SCD only uses one column of  $\widetilde{\mathbf{A}}$  per iteration, that is, one sample, to compute  $\nabla_{i_k}\Phi(\mathbf{u}^k) = (1/(n^2\mu))\widetilde{\mathbf{A}}^T_{i_k}\mathbf{s}^k$ .

Now, we come to the convergence rate of SCD [98], which is described as

$$\mathbb{E}[D(\mathbf{u}^k)] - D(\mathbf{u}^*) \le \min\left\{ \left(1 - \frac{\mu}{n}\right)^k, \frac{n}{n+k} \right\} C \quad (16)$$

in a unified style for strongly convex and nonstrongly convex problems, where  $C = D(\mathbf{u}^0) - D(\mathbf{u}^*) + ||\mathbf{u}^0 - \mathbf{u}^*||_L^2$ .

We can also perform Nesterov's acceleration technique to accelerate SCD by combining it with (6a)–(6c). When  $\Phi(\mathbf{u})$  in (15) is strongly convex, the resultant method is called accelerated randomized proximal coordinate gradient (APCG) [16], and it is described in Algorithm 3. When the strong convexity assumption is absent, the method is called Accelerated Parallel PROXimal (APPROX) coordinate descent [15], and it is written in Algorithm 4, where  $\theta_k > 0$  is obtained by solving equation  $\theta_k^2 = (1 - \theta_k)\theta_{k-1}^2$ , which is initialized as  $\theta_0 = 1/n$ . Especially, APPROX reduces to (6a)–(6c) when n=1. Both APCG and APPROX have a faster convergence rate than SCD, which is given as follows in a unified style:

$$\mathbb{E}[D(\mathbf{u}^k)] - D(\mathbf{u}^*) \le \min\left\{ \left(1 - \frac{\sqrt{\mu}}{n}\right)^k, \left(\frac{2n}{2n+k}\right)^2 \right\} C$$

where C is given in (16).

1) Efficient Implementation: Both APCG and APPROX need to perform full-dimensional vector operations in steps (17a), (17d), (18a), and (18d), which make the per iteration cost higher than that of SCD, where the latter only needs to consider one dimension per iteration. This may cause the overall computational cost of APCG and APPROX higher than that of the full AGD. To avoid such a situation, we can use a change of variables scheme, which is firstly proposed in [99] and then adopted in [15] and [16]. Take APPROX as an example. We only need to introduce an auxiliary variable  $\hat{\mathbf{u}}^k$  initialized at  $\mathbf{0}$  and change the

# Algorithm 3 APCG

Input 
$$\mathbf{u}^0 = \mathbf{z}^0$$
.

for 
$$k = 0, 1, \cdots$$
 do

$$\mathbf{y}^k = \frac{1}{1 + \sqrt{\mu}/n} \left( \mathbf{u}^k + \frac{\sqrt{\mu}}{n} \mathbf{z}^k \right),\tag{17a}$$

Randomly sample  $i_k$  from  $\{1, \ldots, n\}$ 

$$\mathbf{z}_{i_k}^{k+1} = \underset{\sim}{\operatorname{argmin}} \left( \Psi_{i_k}(z) + \langle \nabla_{i_k} \Phi(\mathbf{y}^k), z \rangle \right)$$
 (17b)

$$+\frac{L_{i_k}\sqrt{\mu}}{2}\left\|z-\left(1-\frac{\sqrt{\mu}}{n}\right)\mathbf{z}_{i_k}^k-\frac{\sqrt{\mu}}{n}\mathbf{y}_{i_k}^k\right\|^2\right),$$

$$\mathbf{z}_{i}^{k+1} = \mathbf{z}_{i}^{k}, \forall j \neq i_{k},\tag{17c}$$

$$\mathbf{u}^{k+1} = \mathbf{y}^k + \sqrt{\mu} \left( \mathbf{z}^{k+1} - \mathbf{z}^k \right) + \frac{\mu}{n} \left( \mathbf{z}^k - \mathbf{y}^k \right),$$
(17d)

end for

# Algorithm 4 APPROX

Input 
$$\mathbf{u}^0 = \mathbf{z}^0$$
.

for 
$$k=0,1,\cdots$$
 do

$$\mathbf{y}^k = (1 - \theta_k)\mathbf{u}^k + \theta_k \mathbf{z}^k, \tag{18a}$$

Randomly sample  $i_k$  from  $\{1, \ldots, n\}$ 

$$\mathbf{z}_{i_k}^{k+1} = \operatorname{argmin} \left( \Psi_{i_k}(z) + \langle \nabla_{i_k} \Phi(\mathbf{y}^k), z \rangle \right)$$
 (18b)

$$+\frac{n\theta_k L_{i_k}}{2} \|z - \mathbf{z}_{i_k}^k\|^2$$
,

$$\mathbf{z}_{i}^{k+1} = \mathbf{z}_{i}^{k}, \forall j \neq i_{k}, \tag{18c}$$

$$\mathbf{u}^{k+1} = \mathbf{y}^k + n\theta_k \left( \mathbf{z}^{k+1} - \mathbf{z}^k \right), \tag{18d}$$

#### end for

updates by the following ones at each iteration:

$$\begin{split} \mathbf{z}_{i_k}^{k+1} &= \operatorname*{argmin}_z \left( \Psi_{i_k}(z) + \langle \nabla_{i_k} \Phi(\mathbf{z}^k + \theta_k^2 \hat{\mathbf{u}}^k), z \rangle \right. \\ &\left. + \frac{n\theta_k L_{i_k}}{2} \|z - \mathbf{z}_{i_k}^k\|^2 \right) \\ \hat{\mathbf{u}}_{i_k}^{k+1} &= \hat{\mathbf{u}}_{i_k}^k - \frac{1 - n\theta_k}{\theta_k^2} \left( \mathbf{z}_{i_k}^{k+1} - \mathbf{z}_{i_k}^k \right) \\ \hat{\mathbf{u}}_{i_j}^{k+1} &= \hat{\mathbf{u}}_{i_j}^k, \quad \mathbf{z}_{i_j}^{k+1} = \mathbf{z}_{i_j}^k, \quad \forall j \neq i_k. \end{split}$$

The partial gradient  $\nabla_{i_k}\Phi(\mathbf{z}^k+\theta_k^2\hat{\mathbf{u}}^k)$  can be efficiently computed in a way similar to that of SCD discussed above with almost no more burden.

2) Applications to the Regularized Empirical Risk Minimization: Now, we consider the regularized empirical risk minimization problem, which is a special case of problem (1) and is described as

$$\min_{\mathbf{x} \in \mathbb{R}^p} F(\mathbf{x}) \stackrel{\text{def}}{=} \frac{1}{n} \sum_{i=1}^n g_i(\mathbf{A}_i^T \mathbf{x}) + h(\mathbf{x}).$$
 (19)

We often normalize the columns of A to have unit norm. Motivated by (14), we minimize the negative of the dual of (19) as

$$\min_{\mathbf{u} \in \mathbb{R}^n} D(\mathbf{u}) \stackrel{\text{def}}{=} h^* \left( \frac{\mathbf{A}\mathbf{u}}{n} \right) + \frac{1}{n} \sum_{i=1}^n g_i^*(-\mathbf{u}_i)$$
 (20)

where  $h^*(\mathbf{u}) = \max_{\mathbf{v}} \{ \langle \mathbf{u}, \mathbf{v} \rangle - h(\mathbf{v}) \}$  is the convex conjugate of h. For some applications in machine learning, for example, SVM,  $\nabla_i h^*(\mathbf{Au}/n)$ , and  $\mathrm{Prox}_{q_i^*}(u)$  can be efficiently computed [100] and we can use APCG and APPROX to solve (20). When  $g_i$  is L-smooth and h is  $\mu$ -strongly convex, APCG needs  $O((n + \sqrt{nL/\mu})\log(1/\varepsilon))$ iterations to obtain an  $\varepsilon$ -approximate expected dual gap  $\mathbb{E}[F(\mathbf{x}^k)] + \mathbb{E}[D(\mathbf{u}^k)] \leq \varepsilon$  [16]. When the smoothness assumption on  $g_i$  is absent, the required iteration number of APPROX is  $O(n \log n + \sqrt{n/\epsilon})$  for the expected  $\varepsilon$  dual

One limitation of the SCD-based methods is that they require computing  $\nabla_i h^*(\mathbf{Au}/n)$  and  $\mathrm{Prox}_{g_i^*}(u)$ , rather than  $\operatorname{Prox}_h(\mathbf{x})$  and  $\nabla g_i(\mathbf{A}_i^T\mathbf{x})$ . In some applications, for example, regularized logistic regression, the SCD-based methods need inner loops and they are less efficient than the VR-based methods. However, for other applications where the VR-based methods cannot be used, for example,  $g_i$  is nonsmooth, the SCD-based method may be a better choice, especially when h is chosen as the  $\ell_2$  regularizer and the proximal mapping of  $q_i$  is simple.

3) Restart for SVM Under the Quadratic Growth Condition: In machine learning, some problems may satisfy a condition that is weaker than strong convexity and stronger than convexity, namely the quadratic growth condition [101]. For example, consider the dual problem of SVM [102]. Can we expect a faster convergence than the sublinear rate of O(1/k) or  $O(1/k^2)$ ? The answer is yes. Some studies have shown that the accelerated methods with restart [103]-[105] enjoy a linear convergence under the quadratic growth condition. Generally, if we have an accelerated method with an  $O(1/k^2)$  rate at hand, for example, APPROX, we can run the method without any change and restart it after several iterations with warm starts. If we set the restart period according to the quadratic growth condition constant, a similar constant to the condition number in the strong convexity assumption, the resultant method converges with a linear rate, which is faster than the nonaccelerated counterparts. Moreover, when the quadratic growth condition constant is unknown (it is often the case in practice), Li and Lin [18] and Fercoq and Qu [104], [105] showed that the method also converges linearly, but the rate may not be optimal.

4) Nonuniform Sampling: In problem (14), we have  $L_i = \|\widetilde{\mathbf{A}}_i\|^2/(n^2\mu)$ . For the analysis in Section III-B2, we normalize the columns of  $\mathbf{A}$  to have unit norm and  $L_i$ have the same values for all i. When they are not normalized, a variety of works have focused on the nonuniform sampling of the training sample  $i_k$  [99], [106]–[108]. For example, Allen-Zhu et al. [107] selected the ith sample with probability proportional to  $\sqrt{L_i}$  and obtained better performance than the uniform sampling scheme. Intuitively speaking, when  $L_i$  is large, the function is less smooth along the ith coordinate, so we should sample it more often to balance the overall convergence speed.

# C. Primal-Dual Method and Its Accelerated **Stochastic Variants**

The VR-based methods and SCD-based methods perform in the primal space and the dual space, respectively. In this section, we introduce another common scheme, namely the primal-dual-based methods [46], [47], which perform both in the primal space and the dual space. Consider problem (19). It can be written in the min-max

$$\min_{\mathbf{x} \in \mathbb{R}^p} \max_{\mathbf{u} \in \mathbb{R}^n} \frac{1}{n} \sum_{i=1}^n \left( \left\langle \mathbf{A}_i^T \mathbf{x}, \mathbf{u}_i \right\rangle - g_i^*(\mathbf{u}_i) \right) + h(\mathbf{x}). \tag{21}$$

We first introduce the general primal-dual method with Bregman distance to solve problem (21) [20], which consists of the following steps at each iteration:

$$\hat{\mathbf{x}}^k = \alpha(\mathbf{x}^k - \mathbf{x}^{k-1}) + \mathbf{x}^k, \qquad (22a)$$

$$\mathbf{u}^{k+1} = \underset{\mathbf{u}}{\operatorname{argmax}} \left( \frac{1}{n} \langle \mathbf{A}^T \hat{\mathbf{x}}^k, \mathbf{u} \rangle - \frac{1}{n} \sum_{i=1}^n g_i^*(\mathbf{u}_i) - \tau D(\mathbf{u}, \mathbf{u}^k) \right)$$
(22b)

$$\mathbf{x}^{k+1} = \underset{\mathbf{x}}{\operatorname{argmin}} \left( h(\mathbf{x}) + \left\langle \mathbf{x}, \frac{1}{n} \mathbf{A} \mathbf{u}^{k+1} \right\rangle + \frac{\eta}{2} \|\mathbf{x} - \mathbf{x}^{k}\|^{2} \right)$$
(22c)

for constants  $\alpha$ ,  $\tau$ , and  $\eta$  to be specified later and  $\mathbf{x}^{-1} = \mathbf{x}^{0}$ . The primal-dual method alternately maximizes **u** in the dual space and minimizes x in the primal space.

As explained in Section III, dealing with all the samples at each iteration is time-consuming when n is large, so we want to handle only one sample. Accordingly, we can sample only one  $i_k$  randomly in (22b) at each iteration. The resultant method is described in Algorithm 5, and it reduces to the stochastic primal-dual coordinate (SPDC) method proposed in [19] when we take D(u, v) = $(1/2)(u-v)^2$  as a special case. Combining the initialization  $\mathbf{s}^0 = (1/n)\mathbf{A}\mathbf{u}^0$  and the update rules (23c) and (23e), we know  $\mathbf{s}^k = (1/n)\mathbf{A}\mathbf{u}^k$ .

Similar to APCG, when each  $g_i$  is L-smooth, h is  $\mu$ -strongly convex, and the columns of A are normalized to have unit norm, SPDC needs  $O((n + \sqrt{nL/\mu}) \log(1/\epsilon))$ iterations to find a solution such that  $\mathbb{E}[\|\mathbf{x}^k - \mathbf{x}^*\|^2] \leq \varepsilon$ .

One limitation of SPDC is that it applies only to problems when the proximal mappings of  $g_i^*$  and h can be efficiently computed. In some applications, we want to use  $\nabla g_i$ , rather than  $\text{Prox}_{g_i^*}$ . To remedy this problem, Lan and Zhou [20] creatively used the Bregman distance

# Algorithm 5 SPDC

Input 
$$\mathbf{x}^0 = \mathbf{x}^{-1}$$
,  $\tau = \frac{2}{\sqrt{n\mu L}}$ ,  $\eta = 2\sqrt{n\mu L}$ , and  $\alpha = 1 - \frac{1}{n+2\sqrt{nL/\mu}}$  for  $k = 0, 1, \cdots$  do

$$\begin{split} \hat{\mathbf{x}}^k &= \alpha(\mathbf{x}^k - \mathbf{x}^{k-1}) + \mathbf{x}^k, \\ \mathbf{u}_{i_k}^{k+1} &= \operatorname*{argmax}_u \Big( \langle \mathbf{A}_{i_k}^T \hat{\mathbf{x}}^k, u \rangle - g_{i_k}^*(u) - \tau D(u - \mathbf{u}_{i_k}^k) \Big), \end{split} \tag{23a}$$

$$\mathbf{u}_{j}^{k+1} = \mathbf{u}_{j}^{k}, \forall j \neq i_{k},$$

$$\mathbf{x}^{k+1} = \underset{\mathbf{x}}{\operatorname{argmin}} \left( h(\mathbf{x}) + \langle \mathbf{x}, \mathbf{s}^{k} + (\mathbf{u}_{i_{k}}^{k+1} - \mathbf{u}_{i_{k}}^{k}) \mathbf{A}_{i_{k}} \rangle$$
(23c)

$$+\frac{\eta}{2}\|\mathbf{x} - \mathbf{x}^k\|^2\right),\tag{23d}$$

$$\mathbf{s}^{k+1} = \mathbf{s}^k + \frac{1}{n} (\mathbf{u}_{i_k}^{k+1} - \mathbf{u}_{i_k}^k) \mathbf{A}_{i_k}, \tag{23e}$$

end for

induced by  $g_i^*$  in (23b). Specifically, taking  $\varphi$  in (3) as  $g_{i_k}^*$ , letting  $\mathbf{z}_{i_k}^{k-1} = \hat{\nabla} g_{i_k}^*(\mathbf{u}_{i_k}^k)$  and defining  $z = (\mathbf{A}_{i_k}^T \hat{\mathbf{x}}^k + \tau \mathbf{z}_{i_k}^{k-1})/(1+\tau)$ , step (23b) reduces to

$$\begin{aligned} \mathbf{u}_{i_k}^{k+1} &= \operatorname*{argmax}_{u} \left( \langle \mathbf{A}_{i_k}^T \hat{\mathbf{x}}^k + \tau \hat{\nabla} g_{i_k}^* (\mathbf{u}_{i_k}^k), u \rangle - (1+\tau) g_{i_k}^* (u) \right) \\ &= \operatorname*{argmax}_{u} \left( \langle z, u \rangle - g_{i_k}^* (u) \right) = \nabla g_{i_k}(z). \end{aligned}$$

Then, we have  $z\in\partial g_{i_k}^*(\mathbf{u}_{i_k}^{k+1})$  and denote it as  $\mathbf{z}_{i_k}^k$ . Thus, we can replace steps (23b) and (23c) by the following two steps:

$$\begin{split} \mathbf{z}_j^k &= \begin{cases} \mathbf{A}_j^T \hat{\mathbf{x}}^k + \tau \mathbf{z}_j^{k-1} \\ \mathbf{z}_j^{k-1}, & j \neq i_k \end{cases} \\ \mathbf{u}_j^{k+1} &= \begin{cases} \nabla g_j(\mathbf{z}_j^k), & j = i_k \\ \mathbf{u}_j^k, & j \neq i_k. \end{cases} \end{split}$$

Accordingly, the resultant method, called the randomized primal–dual gradient (RPDG) method [20], is only based on  $\nabla g_j(z)$  and the proximal mapping of  $h(\mathbf{x})$ . To find an  $\varepsilon$ -optimal solution, it needs the same number of iterations as SPDC but each iteration has the same computational cost as the VR-based methods, for example, Katyusha.

1) Relation to Nesterov's AGD: It is interesting to study the relationship between the primal–dual method and Nesterov's accelerated gradient method. Lan and Zhou [20] proved that (22a)–(22c) with  $\tau_k = (1-\theta_k)/\theta_k$ ,  $\eta_k = L\theta_k$ , and  $\alpha_k = \theta_k/\theta_{k-1}$ , appropriate Bregman distance reduces to (6a)–(6c) when solving (19), where we use adaptive parameters in (22a)–(22c). Thus, RPDG can also be seen as an extension of Nesterov's AGD to finite-sum stochastic optimization problems.

2) Nonstrongly Convex Problems: When the strong convexity assumption on  $h(\mathbf{x})$  is absent, Chambolle et al. [76] studied the  $O(1/\sqrt{\varepsilon})$  iteration upper bound for the stochastic primal-dual hybrid gradient algorithm, which is a variant of SPDC. However, no explicit dependence on nwas given in [76]. On the other hand, the perturbation approach is a popular way to obtain sharp convergence results for nonstrongly convex problems. Specifically, define a perturbation problem by adding a small perturbation term  $\varepsilon \|\mathbf{x}^0 - \mathbf{x}\|^2$  to problem (19) and solve it by RPDG, which is developed for strongly convex problems. However, the resultant gradient complexity has an additional term  $\log(1/\varepsilon)$  when compared with the lower bound in [67] and the upper bound in [13]. Since the conditions in the HOOD framework [84] may not be satisfied for RPDG due to the dual term, currently the reduction approach introduced in Section III-A2 has not been applied to the primal-dual-based methods to remove the additional poly-logarithmic factor.

# IV. ACCELERATED NONCONVEX ALGORITHMS

In this section, we introduce the generalization of acceleration to nonconvex problems. Specifically, Section IV-A introduces the deterministic algorithms and Section IV-B the stochastic ones.

# A. Deterministic Algorithms

In Sections IV-A1 and IV-A2, we describe the algorithms to find first-order and second-order stationary points, respectively.

1) Achieving First-Order Stationary Point: GD and its proximal variant are widely used in machine learning, both for convex and nonconvex applications. For nonconvex problems, GD finds an  $\varepsilon$ -approximate first-order stationary point within  $O(1/\varepsilon^2)$  iterations [33].

Motivated by the success of heavy-ball method, Ochs *et al.* [109] studied its nonconvex extension with the name of iPiano. Specifically, consider problem (1) with smooth (possibly nonconvex)  $f(\mathbf{x})$  and convex  $h(\mathbf{x})$  (possibly nonsmooth), and the heavy-ball method (4) with  $\beta \in [0,1)$  and  $\eta < 2(1-\beta)/L$ . Ochs *et al.* [109] proved that any limit point  $\mathbf{x}^*$  of  $\mathbf{x}^k$  is a critical point of (1), that is,  $0 \in \nabla f(\mathbf{x}^*) + \partial h(\mathbf{x}^*)$ . Moreover, the number of iterations to find an  $\varepsilon$ -approximate first-order stationary point is  $O(1/\varepsilon^2)$ .

Besides the heavy-ball method, some researchers studied the nonconvex accelerated gradient method extended from Nesterov's AGD. For example, Ghadimi and Lan [110] studied the following method for problem (1) with convex  $h(\mathbf{x})$ :

$$\mathbf{y}^k = (1 - \theta_k)\mathbf{x}^k + \theta_k\mathbf{z}^k \tag{24a}$$

$$\mathbf{z}^{k+1} = \operatorname{Prox}_{\delta_k h}(\mathbf{z}^k - \delta_k \nabla f(\mathbf{y}^k))$$
 (24b)

$$\mathbf{x}^{k+1} = \operatorname{Prox}_{\sigma_k h}(\mathbf{y}^k - \sigma_k \nabla f(\mathbf{y}^k)) \tag{24c}$$

which is motivated by (6a)–(6c). In fact, when  $h(\mathbf{x}) = 0$ ,  $\delta_k = 1/(L\theta_k)$ , and  $\sigma_k = 1/L$ , (6a)–(6c) and (24a)–(24c) are equivalent. Ghadimi and Lan [110] proved that (24a)–(24c) needs  $O(1/\varepsilon^2)$  iterations to find an  $\varepsilon$ -approximate first-order stationary point by setting  $\theta_k = 2/(k+1)$ ,  $\sigma_k = 1/(2L)$ , and  $\sigma_k \leq \delta_k \leq (1+\theta_k/4)\sigma_k$ . On the other hand, when  $f(\mathbf{x})$  is also convex, (24a)–(24c) have the optimal  $O(1/\sqrt{\varepsilon})$  iteration complexity to find an  $\varepsilon$ -optimal solution by a different setting of  $\delta_k = k\sigma_k/2$ .

Although (24a)-(24c) guarantee the convergence for nonconvex programming while maintaining the acceleration for convex programming, one disadvantage is that the parameter settings for convex and nonconvex problems are different. To address this issue, Li and Lin [111] proposed the following method:

$$\begin{aligned} &\mathbf{y}_k = \mathbf{x}_k + \frac{\theta_k}{\theta_{k-1}}(\mathbf{z}_k - \mathbf{x}_k) + \frac{\theta_k(1 - \theta_{k-1})}{\theta_{k-1}}(\mathbf{x}_k - \mathbf{x}_{k-1}) \\ &\mathbf{z}_{k+1} = \operatorname{Prox}_{\eta h}(\mathbf{y}_k - \eta \nabla f(\mathbf{y}_k)) \\ &\mathbf{v}_{k+1} = \operatorname{Prox}_{\eta h}(\mathbf{x}_k - \eta \nabla f(\mathbf{x}_k)) \\ &\mathbf{x}_{k+1} = \begin{cases} \mathbf{z}_{k+1}, & \text{if } F(\mathbf{z}_{k+1}) \leq F(\mathbf{v}_{k+1}) \\ \mathbf{v}_{k+1}, & \text{otherwise} \end{cases} \end{aligned}$$

which is motivated by the monotone AGD proposed in [112]. Intuitively, the first two steps perform a proximal AGD update with the same update rule of  $\theta_k$  as that in (5a) and (5b), the third step performs a proximal GD update, and the last step chooses the one with the smaller objective. Similar to the heavy-ball method, Beck and Teboulle [112] proved that any limit point of  $\mathbf{x}^k$  is a critical point, and the method needs  $O(1/\varepsilon^2)$  iterations to find an  $\varepsilon$ -approximate first-order stationary point. On the other hand, when both  $f(\mathbf{x})$  and  $h(\mathbf{x})$  are convex, the same  $O(1/\sqrt{\varepsilon})$  iteration complexity as Nesterov's AGD is maintained. Moreover, the algorithm for convex programming and nonconvex programming keeps the same parameters. The price paid is that the computational cost per iteration of the above method is higher than that of (24a)–(24c).

Besides the above algorithms, Nesterov et al. [42] have also extended his AGD to nonconvex. Similar to the GD [48] discussed in Section II-D, Nesterov's method also needs a line search and thus it is not a rigorously "firstorder" method.

We can see that none of the above algorithms have provable improvement after adopting the technique of heavy-ball method or Nesterov's AGD. One may ask: can we find a provable faster accelerated gradient method for nonconvex programming? The answer is yes. Carmon et al. [22] proposed a method which achieves an  $\varepsilon$ -approximate first-order stationary point within  $O(1/\varepsilon^{7/4})$  gradient and function evaluations. The algorithm in [22] is complex to implement, so we omit the details.

2) Achieving Second-Order Stationary Point: We first discuss whether GD can find the approximate second-order stationary point. To answer this question, Jin et al. [113] studied a simple variant of GD with appropriate perturbations and showed that the method achieves an  $O(\varepsilon, O(\sqrt{\varepsilon}))$ -approximate second-order stationary point within  $\widetilde{O}(1/\varepsilon^2)$  iterations, where  $\widetilde{O}$  hides the polylogarithmic factors. We can see that this rate is exactly the rate of GD to first-order stationary point, with only the additional log factor. The method proposed in [113] is given in Algorithm 6, where  $Uniform(B_0(r))$  means the perturbation uniformly sampled from a ball with radius r.

# **Algorithm 6** Perturbed GD

```
\begin{array}{l} \text{Input } \mathbf{x}^0 = \mathbf{z}, \; p = 0, \; T = \widetilde{O}(\frac{1}{\sqrt{\varepsilon}}), \; r = \widetilde{O}(\varepsilon), \; \eta = O(\frac{1}{L}), \\ \text{and } \varepsilon' = O(\varepsilon^{1.5}). \end{array}
for k=0,1,\cdots do
    if \|\nabla f(\mathbf{x}^k)\| \le \varepsilon and k > p + T (i.e., no perturbation
     in last T steps) then
        \mathbf{z} = \mathbf{x}^k, \ p = k

\mathbf{x}^k = \mathbf{x}^k + \boldsymbol{\xi}^k, \quad \boldsymbol{\xi}^k \sim \mathrm{Uniform}(B_0(r)),
    \mathbf{x}^{k+1} = \mathbf{x}^k - \eta \nabla f(\mathbf{x}^k),
    if k = p + T and f(\mathbf{z}) - f(\mathbf{x}^k) < \varepsilon' then
         break,
     end if
end for
```

Intuitively, when the norm of the current gradient is small, it indicates that the current iterate is potentially near a saddle point or a local minimum. If it is near a saddle point, the uniformly distributed perturbation helps to escape it, which is added at most once in every T iterations. On the other hand, when the objective almost does not decrease after T iterations from last perturbation, it achieves the local minimum with high probability and we can stop the algorithm.

Besides [113], Lee et al. [114] showed that the plain GD without perturbations almost always escapes saddle points asymptotically. However, it may take exponential time [115].

Now, we come to the accelerated gradient method. Built upon Algorithm 6 and (5a) and (5b), Jin et al. [24] proposed a variant of AGD with perturbations and showed that the method needs  $O(1/\varepsilon^{7/4})$  iterations to achieve an  $(\varepsilon, O(\sqrt{\varepsilon}))$ -approximate second-order stationary point, which is faster than the perturbed GD. We describe the method in Algorithm 7, where the negative curvature exploitation (NCE) step chooses  $\mathbf{x}^{k+1}$  to be  $\mathbf{x}^k$  +  $\delta$  or  $\mathbf{x}^k - \delta$  whichever having a smaller objective f, where  $\delta = s\mathbf{v}^k/\|\mathbf{v}^k\|$  for some constant s.

In the above scheme, the first "if" step is similar to the perturbation step in the perturbed GD. The following three steps are similar to the AGD steps in (5a) and (5b), where  $\mathbf{v}^k$  is the momentum term in (5a). When the function has large negative curvature between  $\mathbf{x}^k$  and  $\mathbf{y}^k$ , that is, the second "if" condition holds, NCE simply moves along the direction based on the momentum.

# Algorithm 7 Perturbed AGD

Input 
$$\mathbf{x}^0, \mathbf{v}^0, T = \widetilde{O}(\frac{1}{\varepsilon^{1/4}}), r = \widetilde{O}(\varepsilon), \beta = \widetilde{O}(1 - \varepsilon^{1/4}),$$
 $\eta = O(\frac{1}{L}), \gamma = \widetilde{O}(\sqrt{\varepsilon})$  and  $s = \widetilde{O}(\sqrt{\varepsilon}).$ 
for  $k = 0, 1, \cdots$  do

if  $\|\nabla f(\mathbf{x}^k)\| \le \varepsilon$  and no perturbation in last  $T$  steps, then

 $\mathbf{x}^k = \mathbf{x}^k + \boldsymbol{\xi}^k, \quad \boldsymbol{\xi}^k \sim \mathrm{Uniform}(B_0(r)),$ 
end if
 $\mathbf{y}^k = \mathbf{x}^k + \beta \mathbf{v}^k,$ 
 $\mathbf{x}^{k+1} = \mathbf{y}^k - \eta \nabla f(\mathbf{y}^k),$ 
 $\mathbf{v}^{k+1} = \mathbf{x}^{k+1} - \mathbf{x}^k,$ 
if  $f(\mathbf{x}^k) \le f(\mathbf{y}^k) + \langle \nabla f(\mathbf{y}^k), \mathbf{x}^k - \mathbf{y}^k \rangle - \frac{\gamma}{2} \|\mathbf{x}^k - \mathbf{y}^k\|^2$ 
then

 $(\mathbf{x}^{k+1}, \mathbf{v}^{k+1}) = \mathrm{NCE}(\mathbf{x}^k, \mathbf{v}^k),$ 
end if
end for

**Table 3** Iteration Complexity Comparisons Between GD and AGD for Nonconvex Problems. We Hide the Poly-Logarithmic Factors in  $\tilde{O}$ . We Also Hide n Since We Only Consider Deterministic Optimization

First-order Stationary Point		Second-order Stationary Point	
Methods	Iteration Complexity	Methods	Iteration Complexity
GD [33]	$O(1/\varepsilon^2)$	Perturbed GD	$\widetilde{O}(1/\varepsilon^2)$
		[113]	
AGD [22]	$\widetilde{O}(1/\varepsilon^{7/4})$	AGD	$\tilde{O}(1/\varepsilon^{7/4})$
		[21], [23], [24]	

Besides [24], Carmon *et al.* [21] and Agarwal *et al.* [23] also established the  $O(1/\varepsilon^{7/4})$  gradient complexity to achieve an  $\varepsilon$ -approximate second-order stationary point. Carmon *et al.* [21] employed a combination of (regularized) AGD and the Lanczos method, and Agarwal *et al.* [23] proposed a careful implementation of the Nesterov–Polyak method, using accelerated methods for fast approximate matrix inversion.

At last, we compare the iteration complexity of the accelerated methods and nonaccelerated methods in Table 3, including both the approximation of first-order stationary point and second-order stationary point.

# **B. Stochastic Algorithms**

Due to the success of deep neural network, in recent years, people are interested in stochastic algorithms for nonconvex problem (1) or (10) with huge n, especially the accelerated variants. Sutskever  $et\ al.\ [116]$  empirically observed that the following plain stochastic AGD performs well when training deep neural networks:

$$\mathbf{y}^k = \mathbf{x}^k + \beta_k (\mathbf{x}^k - \mathbf{x}^{k-1})$$
$$\mathbf{x}^{k+1} = \mathbf{y}^k - \eta \nabla f_{i_k}(\mathbf{y}^k)$$

where  $\beta_k$  is empirically set as

$$\beta_k = \min\{1 - 2^{-1 - \log_2(\lfloor k/250 \rfloor + 1)}, \beta_{\max}\}$$

and  $\beta_{\text{max}}$  is often chosen as 0.999 or 0.995.

In this section, we introduce the stochastic nonconvex algorithms with more theory supports than the above plain stochastic AGD. For simplicity, we consider problem (10) with each  $f_i(\mathbf{x})$  being L-smooth.

1) Achieving First-Order Stationary Point: When we assume that the variance of the gradient is finite, SGD requires the gradient complexity of  $O(\varepsilon^{-4})$  to achieve an  $\varepsilon$ -approximate first-order stationary point [33]. Similar to stochastic convex optimization, this bound can be further improved by VR. In fact, a slight variant of the SVRG algorithm [117]-[119] and also SAGA [120] achieve the gradient complexity of  $O((n+n^{2/3}\varepsilon^{-2})\wedge\varepsilon^{-10/3})$ , where  $a\wedge$  $b = \min(a, b)$ . This result means that when  $n \to \infty$ , the VR technique can still guarantee a faster convergence rate in the nonconvex stochastic optimization, where we refer this case as the online optimization. However, this bound is still not optimal and it can be further reduced to O((n + $n^{1/2}\varepsilon^{-2}$ )  $\wedge \varepsilon^{-3}$ ) by performing recursive VR [26], [78], [121]–[123]. We take the stochastic path-integrated differential estimator (SPIDER) [26] algorithm as an example. SPIDER can be used for both the finite-sum problem (10) and the online problem. For simplicity, we consider the following simplified method for the finite-sum problem, as shown in Algorithm 8.

# Algorithm 8 SPIDER

$$\begin{aligned} & \mathbf{for} \ k = 0 \ \text{to} \ K \ \mathbf{do} \\ & \mathbf{v}^k = \begin{cases} \nabla f(\mathbf{x}^k), & \text{if } \mathrm{mod} \left(k, n\right) = 0, \\ \nabla f_{i_k}(\mathbf{x}^k) - \nabla f_{i_k}(\mathbf{x}^{k-1}) + \mathbf{v}^{k-1}, & \text{otherwise.} \end{cases} \\ & \eta_k = \min\left(\frac{\varepsilon}{L\sqrt{n}\|\mathbf{v}^k\|}, \frac{1}{2L\sqrt{n}}\right), \\ & \mathbf{x}^{k+1} = \mathbf{x}^k - \eta_k \mathbf{v}^k. \end{aligned}$$

SPIDER is motivated by SVRG, but using a different VR technique. We can compare SPIDER with the loopless SVRG (11a) and (11c) to be more intuitive. SPIDER takes steps along the direction based on past accumulated stochastic gradient information, that is,

$$\mathbf{v}^k = \sum_{t=k_0+1}^k (
abla f_{i_t}(\mathbf{x}^t) - 
abla f_{i_t}(\mathbf{x}^{t-1})) + \mathbf{v}^{k_0}$$

for the latest  $k_0$  such that  $\mod(k_0,n)=0$  and  $\mathbf{v}^{k_0}=\nabla f(\mathbf{x}^{k_0})$ . In contrast, SVRG takes steps only based on the information of current stochastic gradient and the snapshot vector, that is,  $\mathbf{v}^k=\nabla f_{i_k}(\mathbf{x}^k)-\nabla f_{i_k}(\tilde{\mathbf{x}}^s)+\nabla f(\tilde{\mathbf{x}}^s)$ . It was shown in [26] that the variance of  $\mathbf{v}^k$  is smaller than that in the SVRG algorithm by order when  $\mathbf{x}^k$  moves slowly, which contributes to a provably faster convergence rate.

The recursive VR technique was first proposed in the algorithm named SARAH in [78], which was designed for convex optimization and then was extended to nonconvex optimization [123]. For the finite-sum smooth optimization, SPIDER and SARAH almost have the same

	Туре	Algorithm		Online	Finite-Sum
First-order Stationary Point	Original	SGD / GD	[33]	$O(\varepsilon^{-4})$	$O(n\varepsilon^{-2})$
	Acceleration	SVRG / SCSG /SAGA	[117]–[120]	$O\left(\varepsilon^{-10/3}\right)$	$O\left(n+n^{2/3}\varepsilon^{-2}\right)$
		SPIDER / SpiderBoost / SARAH	[26], [121]–[123]	$O\left(\varepsilon^{-3}\right)$	$O\left(n+n^{1/2}\varepsilon^{-2}\right)$
Second-order Stationary Point	Original	Perturbed GD / SGD	[125] [126] [113], [127] [128]	$ \begin{array}{ c c } \widetilde{O}\left(poly(d)\varepsilon^{-4}\right) \\ \widetilde{O}\left(\varepsilon^{-10}\right) \\ \widetilde{O}\left(\varepsilon^{-4}\right) \\ \widetilde{O}\left(\varepsilon^{-3.5}\right) \end{array} $	not given not given $\widetilde{O}\left(n\varepsilon^{-2}\right)$ not given
		NEON+GD/SGD	[129], [130]	$\tilde{O}(\varepsilon^{-4})$	$\widetilde{O}(n\varepsilon^{-2})$
	Acceleration	Perturbed AGD	[24]	not given	$n\hat{\epsilon}^{-1.75}$
(Hessian-smooth Required)		NEON+VR	[25], [117]–[119]	$\widetilde{O}\left(\varepsilon^{-3.5}\right)$	$\widetilde{O}\left(n\varepsilon^{-1.5} + n^{2/3}\varepsilon^{-2}\right)$
• ′		NEON+VR+Momentum	[21], [23], [27]	$\widetilde{O}(\varepsilon^{-3.5})$	$\widetilde{O}\left(n\varepsilon^{-1.5} + n^{3/4}\varepsilon^{-1.75}\right)$

[26]

Table 4 Gradient Complexity Comparisons Between Different Accelerated Stochastic Algorithms and Their Nonaccelerated Counterparts to Find an  $\varepsilon$ -Approximate First-Order Stationary Point or an  $(\varepsilon, O(\varepsilon^{0.5}))$ -Approximate Second-Order Stationary Point for Nonconvex Problems

algorithm form. They are different in the step size. SARAH uses  $\eta = O(1/(L\sqrt{n}))$ , while SPIDER uses a normalized but more conservative step size [at the order of  $O(\varepsilon)$ ]. After [26], some improved versions, such as SpiderBoost [122], also considered allowing a larger step size.

NEON+SPIDER

As for the lower bounds, Fang et al. [26] proved that the gradient complexity of  $O(n + n^{1/2} \varepsilon^{-2})$  matches the lower bound under certain conditions. More recently, Arjevani et al. [124] showed that  $O(\varepsilon^{-3})$  also matches the lower bound when  $n \to \infty$ .

2) Achieving Second-Order Stationary Point: When the objective function is assumed to have a Lipschitz continuous Hessian matrix, acceleration has also been done to find an approximate second-order stationary point. For example, Agarwal et al. [23] and Tripuraneni et al. [27] converted the cubic regularization method [131] for finding a second-order stationary point using stochasticgradient-based and Hessian-vector-product-based methods. Xu et al. [129] and Allen-Zhu and Li [130] proposed a generic saddle point-escaping method called NEON, which approximates Hessian-vector product by stochastic gradient. For the convergence rate, to search an  $(\varepsilon, O(\varepsilon^{0.5}))$ approximate second-order stationary point, in the finitesum case, the VR and the momentum techniques [21], [23] can reduce the gradient complexity to  $O(n\varepsilon^{-1.5} +$  $n^{3/4} \varepsilon^{-1.75}$ ). In the online case, Ge et al. [125] first proved that noisy SGD escapes from saddle points in polynomial times. Later, Daneshmand et al. [126] obtained a gradient complexity of  $\widetilde{O}(\varepsilon^{-10})$ . This bound was finally improved by Fang et al. [128], in which the authors proved that noisy SGD can actually find a second-order stationary point within the gradient complexity of  $\widetilde{O}(\varepsilon^{-3.5})$ . For the

variants of SGD, by fusing negative curvature search with VR, for finding an  $(\varepsilon, O(\varepsilon^{0.25}))$ -approximate second-order stationary point, Allen-Zhu [25] obtained a lower gradient complexity of  $\widetilde{O}(\varepsilon^{-3.25})$ . When using the SPIDER [26] technique, one can obtain a complexity of  $\widetilde{O}(\varepsilon^{-3})$  to find an  $(\varepsilon, O(\varepsilon^{0.5}))$ -approximate second-order stationary point. Table 4 summarizes the gradient complexity comparisons of the existing algorithms.

# V. DISCUSSION AND LIMITATION

Accelerated algorithms have been widely used in machine learning due to their provably faster convergence and simplicity in implementation. In this article, we review the accelerated deterministic algorithms, accelerated stochastic algorithms, and accelerated nonconvex algorithms for machine learning. Due to space limit, our review is incomplete as we have left some interesting topics out, for example, the acceleration for distributed optimization [132]-[142]. Its challenge over the nondistributed algorithms introduced in this article is that we should pay attention to the agreement among different nodes. Modifications are required to extend the classical AGD to distributed optimization.

The efficiency of accelerated algorithms has been verified in practice for convex optimization, either deterministic or stochastic. However, in reality, some complex accelerated nonconvex algorithms seem less efficient. One remarkable example is that although they are proved to converge faster to first-order or second-order stationary points than SGD, when training deep neural networks, they still cannot beat SGD or the plain stochastic AGD. There is still a gap between theory and practice for accelerated nonconvex optimization.

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#### Li et al.: Accelerated First-Order Optimization Algorithms for Machine Learning

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