To Reviewer #1

Q1: “I am not sure if this is significant enough...The improved analysis is useful but not provide a big impact”
A1: Improving the over-parameterization condition from $\Omega(n^{24})$ to $\tilde{\Omega}(n^8)$ without making any additional assumption on the training data is by no means trivial. More importantly, our work clearly points out the right direction to improve the over-parameterization condition: (1) proving tighter gradient lower bound, and (2) proving shorter trajectory length for (stochastic) gradient descent. Our work sheds light on further improving the over-parameterization condition and trigger a lot of followup work to make the over-parameterization well-aligned with the neural network width used in practice. Moreover, we believe our proof idea can also be generalized to studying a broader class of neural networks (e.g., CNN, ResNet). Therefore, we believe our contribution to this line of research is significant and our work has a big impact in the field of deep learning theory.

Q2: “...very different ideas may be needed, I do not see how the new insight can help improve the requirement...”
A2: We don’t fully agree with your comment on this point. Based on our current ideas, there is still a big room to improve the over-parameterization requirement. We explain it as follows: in our paper, the proved over-parameterization condition is $\tilde{\Omega}(n^8)$ when the gradient lower bound is roughly in the order of $O(m/n^2)$ (Here the other problem-dependent parameters are omitted, please refer to Lemma 4.1 for details). If we can improve the gradient lower bound to $O(m/n)$ or even $O(m)$ (in the same order as the gradient upper bound), the over-parameterization condition can be further improved to be $\Omega(n^3)$ or even $\Omega(n^2)$ respectively. This clearly shows the potential and promise of proving tighter gradient lower bound. To give you a more concrete idea, in the current paper, each “gradient region” is designed based on the minimum separation distance $\delta$ and has the same size. However, for high-dimension training data, the separation distance for some data points can be very large, which implies that some “gradient region” can actually has much larger size. Therefore, the total size of “gradient regions” can be greatly enlarged if the average-case separation distance rather than only the smallest one of all training data is taken into consideration. This can potentially lead to larger gradient lower bound. We will point it out in the future work section.

To Reviewer #2

Q1: If the analysis for the case where the top layer weights are also updates.
A1: Thanks for your positive and constructive comments on our paper. In our current submission, we did not optimize the top layer weights since we aim to make a fair comparison between our results and those in existing work (which don’t optimize the top layer). However, we would like to emphasize that our analysis can be easily extended to the case when the top layer is optimized, and the resulting theoretical guarantees (over-parameterization condition and convergence rate) are at least the same as our results. The proof sketch is as follows: similar to our current proof, we can also define a small perturbation region around the initialization, but the new definition involves a constraint on the top layer weights. Then, it can be shown that the neural network enjoys good properties inside such region. Based on such good properties, we can prove that until convergence the neural network weights, including the top layer weights, would not escape from such region. Note that optimizing more parameter can lead to larger gradient, thus we can prove a larger gradient lower bound during the training process which can potential speed up the convergence of optimization algorithm (e.g., GD, SGD). Combining the above analyses, we can derive the corresponding over-parameterization condition and convergence rate, and note that these guarantees would be no worse than existing results presented in our paper. We will briefly discuss this extension in the final version.

To Reviewer #3

Q1: “The complexity increase ... it is not apparent if this treatment was done justly under 4.1.”
A1: We apologize that we did not make it clear, but this is a misunderstanding on the exploitation of the gradient regions. We apologize that we did not make it clear, but this is a misunderstanding on the exploitation of the gradient regions. We would like to clarify that exploiting gradient regions for all the data points is in the proof level, and has nothing to do with the algorithm. Therefore, it would never cause additional computation complexity for the algorithm (e.g., GD and SGD), and it will not affect the complexity calculations in our paper. In other words, the complexity calculations in our current paper are correct.

Q2: “ The remark under 3.11 sounds like it could be better explained for the case of 2 layer ReLU networks.”
A2: We are sorry for making you confused about Remark 3.11. In Remark 3.11, we would like to say that for 2-layer ReLU networks, we can derive its result from Theorem 3.8, because the result of Theorem 3.8 is for any depth $L \geq 2$, and therefore it is also applicable to 2-layer ReLU networks when choosing $L = 2$. However, we can prove stronger results specifically for 2-layer ReLU networks, by exploiting the special structure of 2-layer ReLU networks. Such stronger results are stated in Theorem 3.10. We use Remark 3.11 to compare the results in Theorem 3.8 when choosing $L = 2$, with the results in Theorem 3.11, and show what improvement we can achieve in Theorem 3.11. We hope this clarify your confusion.

Q3: “It would also been great to include practical results for implementation ...”
A3: Thank you for your suggestion. We will consider adding the experimental results in the appendix in the final version.