# Big Bird: Transformers for Longer Sequences

Manzil Zaheer Guru Guruganesh MANZILZ@GOOGLE.COM GURUG@GOOGLE.COM

AVINAVADUBEY@GOOGLE.COM

Google Research, Mountain View, CA, USA

Avinava Dubey Joshua Ainslie, Chris Alberti, Santiago Ontanon, Philip Pham, Anirudh Ravula, Qifan Wang, Li Yang, Amr Ahmed

Google Research, USA

## Abstract

Transformers-based models, such as BERT, have been one of the most successful deep learning models for NLP. Unfortunately, one of their core limitations is the quadratic dependency (mainly in terms of memory) on the sequence length due to their full attention mechanism. To remedy this, we propose, BIGBIRD, a sparse attention mechanism that reduces this quadratic dependency to linear. We show that BIGBIRD is a universal approximator of sequence functions and is Turing complete, thereby preserving these properties of the quadratic, full attention model. Along the way, our theoretical analysis reveals some of the benefits of having O(1) global tokens (such as CLS), that attend to the entire sequence as part of the sparse attention mechanism. The proposed sparse attention can handle sequences of length up to 8x of what was previously possible using similar hardware. As a consequence of the capability to handle longer context, BIGBIRD drastically improves performance on various NLP tasks such as question answering and summarization. We also propose novel applications to genomics data.

# 1. Introduction

Models based on Transformers [92], such as BERT [23, 64], are wildly successful for a wide variety of Natural Language Processing (NLP) tasks and consequently are mainstay of modern NLP research. Their versatility and robustness are the primary drivers behind the wide-scale adoption of Transformers. The model is easily adapted for a diverse range of sequence based tasks – as a seq2seq model for translation [92], summarization [67]. generation [16], etc. or as a standalone encoders for sentiment analysis [84], POS tagging [66], machine reading comprehension [94], etc. – and it is known to vastly outperform previous sequence models like LSTM [38]. The key innovation in Transformers is the introduction of a self-attention mechanism, which can be evaluated in parallel for each token of the input sequence, eliminating the sequential dependency in recurrent neural networks, like LSTM. This parallelism enables Transformers to leverage the full power of modern SIMD hardware accelerators like GPUs/TPUs, thereby facilitating training of NLP models on datasets of unprecedented size. This ability to train on large scale data has led to surfacing of models like BERT [23] and T5 [76], which pretrain transformers on large general purpose corpora and transfer the knowledge to down-stream task. The pretraining has led to significant improvement in low data downstream tasks [52] as well as tasks with sufficient data [102] and thus have been a major force behind the ubiquity of transformers in contemporary NLP. The self-attention mechanism overcomes constraints of RNNs (namely the sequential nature of RNN) by allowing each token in the input sequence to attend independently to every other token in the sequence. This design choice has several interesting repercussions. In particular, the full self-attention have computational and memory requirement that is quadratic in the sequence length. Using commonly available current hardware and model sizes, this requirement translates to roughly being able to handle input sequences of length 512 tokens. This reduces its direct applicability to tasks that require larger context, like question answering [61], document summarization [21], etc.

However, while we know that self-attention and Transformers are useful, our theoretical understanding is rudimentary. What aspects of the self-attention model are necessary for its performance? What can we say about the expressivity of Transformers and similar models? Apriori, it was not even clear from the design if the proposed self-attention mechanism was as effective as RNNs. For example, the self-attention does not even obey sequence order as it is permutation equivariant. This concern has been partially resolved, as Yun et al. [105] showed that transformers are expressive enough to capture all continuous sequence to sequence functions with a compact domain. Meanwhile, Pérez et al. [73] showed that the full transformer is Turing Complete (i.e. can simulate a full Turing machine). Two natural questions arise: Can we achieve the empirical benefits of a fully quadratic self-attention scheme using fewer inner-products? Do these sparse attention mechanisms preserve the expressivity and flexibility of the original network?

In this paper, we address both the above questions and produce a sparse attention mechanism that improves performance on a multitude of tasks that require long contexts. We systematically develop BIGBIRD, an attention mechanism whose complexity is linear in the number of tokens (Sec. 2). We take inspiration from graph sparsification methods and understand where the proof for expressiveness of Transformers breaks down when full-attention is relaxed to form the proposed attention pattern. This understanding helped us develop BIGBIRD, which is theoretically as expressive and also empirically useful. In particular, our BIGBIRD consists of three main part:

- A set of g global tokens that attend on all parts of the sequence.
- For each query  $q_i$ , a set of r random keys that each query will attend to.
- A block of local neighbors w so that each node attends on their local structure.

Our design leads to empirically high performing attention mechanism scaling to much longer sequence lengths (8x) on standard hardware ( $\sim$ 16GB memory) for large size models. Use of gradient checkpointing [15] could potentially allow for handling even longer sequences.

To summarize, our main contributions are:

- 1. BigBird satisfies all the known theoretical properties of full transformer (Sec. 3). In particular, we show that adding extra tokens allows one to express all continuous sequence to sequence functions with only O(n)-inner products. Furthermore, we show that under standard assumptions regarding precision, BigBird is Turing complete.
- 2. Empirically, we show that the extended context modelled by BigBird greatly benefits variety of NLP tasks. In particular, we achieve *state of the art* results for question answering and document summarization on a number of different datasets. Summary of these results are presented in Sec. 4.

3. Lastly, we introduce a novel application of attention based models where long contexts are beneficial: extracting contextual representations of genomics sequences like DNA. With longer masked LM pretraining, Bigbird improves performance on downstream tasks such as promoter-region and chromatin profile prediction (Sec. 5).

## 1.1 Related Work

There have been a number of interesting attempts, that were aimed at alleviating the quadratic dependency of Transformers, which can broadly categorized into two directions. First line of work embraces the length limitation and develops method around it. Simplest methods in this category just employ sliding window [94], but in general most work fits in the following general paradigm: using some other mechanism select a smaller subset of relevant contexts to feed in the transformer and optionally iterate, i.e. call transformer block multiple time with different contexts each time. Most prominently, SpanBERT [43], ORQA [55], REALM [35], RAG [58] have achieved strong performance for different tasks. However, it is worth noting that these methods often require significant engineering efforts (like back prop through large scale nearest neighbor search) and are hard to train.

Second line of work questions if full attention is essential and have tried to come up with approaches that do not require full attention, thereby reducing the memory and computation requirements. Prominently, Dai et al. [22], Sukhbaatar et al. [83], Rae et al. [75] have proposed auto-regresive models that work well for left-to-right language modeling but suffer in tasks which require bidirectional context. Child et al. [17] proposed a sparse model that reduces the complexity to  $O(N\sqrt{N})$ , Kitaev et al. [50] further reduced the complexity to  $O(N \log(N))$  by using LSH to compute nearest neighbors. Ye et al. [104] proposed binary partitions of the data where as Qiu et al. [74] reduced complexity by using block sparsity. Recently, Longformer [8], was introduced that uses a localized sliding window based mask to reduce computation and extended BERT to longer sequence based tasks. Finally, our work is closely related to and built on the work of Extended Transformers Construction [4]. This work was designed to encode structure in text for transformers. The idea of global tokens was used extensively by them to achieve their goals. Our theoretical work can be seen as providing a justification for the success of these models as well. However, it is important to note that most of the aforementioned methods are heuristic based and empirically are not as versatile and robust as the original transformer, i.e. the same architecture do not attain SoTA on multiple standard benchmarks nor handle both encoding and decoding. Moreover, these approximations do not come with theoretical guarantees.

# 2. BigBird Architecture

In this section, we describe the BIGBIRD model using the generalised attention mechanism that is used in each layer of transformer operating on an input sequence  $\mathbf{X} = (\mathbf{x}_1, ..., \mathbf{x}_n) \in \mathbb{R}^{n \times d}$ . The generalized attention mechanism is described by a directed graph D whose vertex set is  $[n] = \{1, ..., n\}$ . The set of arcs (directed edges) represent the set of inner products that the attention mechanism will consider. Let N(i) denote the out-neighbors set of node i in D, then the i<sup>th</sup> output vector of the generalized attention mechanism is defined as

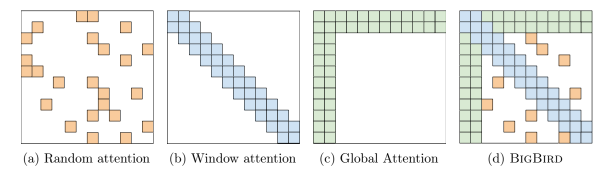


Figure 1: Building blocks of the attention mechanism used in BIGBIRD. White color indicates absence of attention. (a) random attention with r = 2, (b) sliding window attention with w = 3 (c) global attention with g = 2. (d) the combined BIGBIRD model.

$$ATTN_D(\boldsymbol{X})_i = \boldsymbol{x}_i + \sum_{h=1}^{H} \sigma\left(Q_h(\boldsymbol{x}_i) K_h(\boldsymbol{X}_{N(i)})^T\right) \cdot V_h(\boldsymbol{X}_{N(i)})$$
(AT)

where  $Q_h, K_h : \mathbb{R}^d \to \mathbb{R}^m$  are query and key functions respectively,  $V_h : \mathbb{R}^d \to \mathbb{R}^d$  is a value function,  $\sigma$  is a scoring function (e.g. softmax or hardmax) and H denotes the number of heads. Also note  $X_{N(i)}$  corresponds to the matrix formed by only stacking  $\{x_j : j \in N(i)\}$  and not all the inputs.

If D is the complete digraph, we recover the full quadratic attention mechanism of Vaswani et al. [92]. To simplify our exposition, we will operate on the adjacency matrix A of the graph D even though the underlying graph maybe sparse. To elaborate,  $A \in [0,1]^{n \times n}$  with A(i,j)=1 if query i attends to key j and is zero otherwise. For example, when A is the ones matrix (as in BERT), it leads to quadratic complexity, since all tokens attend on every other token. This view of self-attention as a fully connected graph allows us to exploit existing graph theory to help reduce its complexity. The problem of reducing the quadratic complexity of self-attention can now be seen as a graph sparsification problem. It is well-known that random graphs are expanders and can approximate complete graphs in a number of different contexts including in their spectral properties [81, 39]. We believe sparse random graph for attention mechanism should have two desiderata: small average path length between nodes and a notion of locality, each of which we discuss below.

Let us consider the simplest random graph construction, known as Erdős-Rényi model, where each edge is independently chosen with a fixed probability. In such a random graph with just  $\tilde{\Theta}(n)$  edges, the shortest path between any two nodes is logarithmic in the number of nodes [18, 44]. As a consequence, such a random graph approximates the complete graph spectrally and its second eigenvalue (of the adjacency matrix) is quite far from the first eigenvalue [9, 10, 6]. This property leads to a rapid mixing time for random walks in the graph, which informally suggests that information can flow fast between any pair of nodes. Thus, we propose a sparse attention where each query attends over r random number of keys. Mathematically this means that  $A(i,\cdot) = 1$  for r randomly chosen keys (see Fig. 1a).

The second viewpoint which inspired the creation of BigBird is that in most contexts within NLP and computational biology have data which displays a great deal of *locality of reference*. In this phenomenon, a great deal of information about a token can be derived from its neighboring tokens. Most pertinently, Clark et al. [20] investigated self-attention

models in NLP tasks and concluded that that neighboring inner-products are extremely important. The concept of locality, proximity of tokens in linguistic structure, also forms the basis of various linguistic theories such as transformational-generative grammar. In the terminology of graph theory, clustering coefficient is a measure of locality of connectivity. and is high when the graph contains many cliques or near-cliques (subgraphs that are almost fully interconnected). Simple Erdos-Renyi random graphs do not have a high clustering coefficient [85], but a class of random graphs, known as small world graphs, exhibit high clustering coefficient [95]. A particular model introduced by Watts and Strogatz [95] is of high relevance to us as it achieves a good balance between average shortest path and the notion of locality. The generative process of their model is as follows: Construct a regular ring lattice, a graph with n nodes each connected to w neighbors, w/2 on each side. In other words we begin with a sliding window on the nodes. Then a random subset (k%) of all connections is replaced with a random connection. The other (100 - k)% local connections are retained. However, deleting such random edges might be inefficient on modern hardware, so we retain it, which will not affect its properties. In summary, to capture these local structures in the context, in BIGBIRD, we define a sliding window attention, so that during self attention of width w, query at location i attends from i - w/2 to i + w/2 keys. In our notation, A(i, i - w/2 : i + w/2) = 1 (see Fig. 1b).

As an initial sanity check, we performed basic experiments to test whether these intuitions are sufficient in getting performance close to BERT like models, while keeping attention linear in the number of tokens. We found that random blocks and local window were insufficient in capturing all the context necessary to compete with the performance of BERT as seen in Tab. 1.

Model	MLM	SQuAD	MNLI
BERT-base	64.2	88.5	83.4
Random (R)	60.1	83.0	80.2
Window (W)	58.3	76.4	73.1
R + W	62.7	85.1	80.5
Global+R+W	64.4	87.2	82.9

Table 1: Building block comparison @512

The final piece of BIGBIRD is inspired from our theoretical analysis (Sec. 3), which is critical for empirical performance. More specifically, our theory utilizes the importance of "global tokens" (tokens that attend to all tokens in the sequence and to whom all tokens attend to (see Fig. 1c). These global tokens can be defined in two ways:

- BIGBIRD-ITC: In internal transformer construction (ITC), we make some existing tokens "global", which attend over the entire sequence. Concretely, we choose a subset G of indices (with g := |G|), such that A(i, :) = 1 and A(:, i) = 1 for all  $i \in G$ .
- BIGBIRD-ETC: In extended transformer construction (ETC), we include additional "global" tokens such as CLS. Concretely, we add g global tokens that attend to all existing tokens. In our notation, this corresponds to creating a new matrix  $B \in [0,1]^{(N+g)\times(N+g)}$  by adding g rows to matrix A, such that B(i,:)=1, and B(:,i)=1 for all  $i\in\{1,2,\ldots g\}$ , and  $B(g+i,g+j)=A(i,j)\forall\ i,j\in\{1,\ldots,N\}$ . This adds extra location to store context and as we will see in the experiments improves performance.

The final attention mechanism for BigBird (Fig. 1d) has all three of these properties: queries attend to r random keys, each query attends to w/2 tokens to the left of its location and w/2 to the right of its location and they contain g global tokens (The global tokens can be from existing tokens or extra added tokens). We provide implementation details in App. D.

# 3. Theoretical Results about Sparse Attention Mechanism

In this section, we will show that that sparse attention mechanisms are as powerful and expressive as full-attention mechanisms in two respects. First, we show that when sparse attention mechanisms are used in a standalone encoder (such as BERT), they are Universal Approximators of sequence to sequence functions in the style of Yun et al. [105]. Second, we will show that sparse encoder-decoder transformers are Turning Complete (assuming the same conditions defined in [73]).

Complementing the above positive results, we also show that moving to a sparse-attention mechanism incurs a cost, i.e. there is no free lunch. In Sec. 3.4, we show lower bounds by exhibiting a natural task where any sufficiently sparse mechanism will require polynomially more layers.

# 3.1 Notation

The complete Transformer encoder stack is nothing but the repeated application of a single-layer encoder (with independent parameters). We denote the class of such Transformer encoders stack, defined using the generalized encoder (Sec. 2), by  $\mathcal{T}_D^{H,m,q}$  which consists of H-heads with head size m and q is the hidden layer size of the output network, and the attention layer is defined by the directed graph D.

The key difference between our proposed attention mechanism to that of Vaswani et al. [92], Yun et al. [105] is that we add a special token at the beginning of each sequence and assign it a special vector. We will refer to this as  $x_0$ . Therefore our graph D will have vertex set  $\{0\} \cup [n] = \{0, 1, 2, ..., n\}$ . We will assume that this extra node and its respective vector will be dropped at the final output layer of transformer. To avoid cumbersome notation, we will still treat transformer as mapping sequences  $X \in \mathbb{R}^{n \times d}$  to  $\mathbb{R}^{n \times d}$ . We will also allow the transformer to append position embeddings  $E \in \mathbb{R}^{d \times n}$  to matrix X in the input layer.

Finally, we need to define the function class and distance measure for proving universal approximation property. Let  $\mathcal{F}_{CD}$  denote the set of continuous functions  $f:[0,1]^{n\times d}\to\mathbb{R}^{n\times d}$  which are continuous with respect to the topology defined by  $\ell_p$  norm. Recall for any  $p\geq 1$ , the  $\ell_p$  distance is  $d_p(f_1,f_2)=\left(\int \|f_1(X)-f_2(X)\|_p^p dX\right)^{1/p}$ .

## 3.2 Universal Approximators

**Definition 1.** The star-graph S centered at 0 is the graph defined on  $\{0, \ldots, n\}$ . The neighborhood of all vertices i is  $N(i) = \{0, i\}$  for  $i \in \{1, \ldots, n\}$  and  $N(0) = \{1, \ldots, n\}$ .

Our main theorem is that the sparse attention mechanism defined by any graph containing S is a universal approximator:

**Theorem 1.** Given  $1 and <math>\epsilon > 0$ , for any  $f \in \mathcal{F}_{CD}$ , there exists a transformer with sparse-attention,  $g \in \mathcal{T}_D^{H,m,q}$  such that  $d_p(f,g) \leq \epsilon$  where D is any graph containing star graph S.

To prove the theorem, we will follow the standard proof structure outlined in [105].

Step 1: Approximate  $\mathcal{F}_{CD}$  by piece-wise constant functions. Since f is a continuous function with bounded domain  $[0,1)^{n\times d}$ , we will approximate it with a suitable piece-wise constant function. This is accomplished by a suitable partition of the region [0,1) into a grid of granularity  $\delta$  to get a discrete set  $\mathbb{G}_{\delta}$ . Therefore, we can assume that we are dealing with a function  $\bar{f}: \mathbb{G}_{\delta} \to \mathbb{R}^{n\times d}$ , where  $d_p(f,\bar{f}) \leq \frac{\epsilon}{3}$ .

Step 2: Approximate piece-wise constant functions by modified transformers. This is the key step of the proof where the self-attention mechanism is used to generate a contextual-mapping of the input. Informally, a contextual mapping is a unique code for the pair consisting of a matrix  $(X, x_i)$  and a column. Its uniqueness allows the Feed forward layers to use each code to map it to a unique output column.

The main technical challenge is computing the contextual mapping using only sparse attention mechanism. This was done in [105] using a "selective" shift operator which shift up entries that are in a specific interval. Key to their proof was the fact that the shift, was exactly the range of the largest entry to the smallest entry.

Creating a contextual mapping with a sparse attention mechanism is quite a challenge. In particular, because each query only attends to a few keys, it is not at all clear that sufficient information can be corralled to make a contextual embedding of the entire matrix. To get around this, we develop a sparse shift operator which shifts the entries of the matrices if they lie in a certain range. The exact amount of the shift is controlled by the directed sparse attention graphy D. The second key ingredient is the use of additional global token. By carefully applying the operator to a set of chosen ranges, we will show that each column will contain a unique mapping of the full mapping. Therefore, we can augment the loss of inner-products in the self attention mechanism by using multiple layers and an auxiliary global token.

Step 3: Approximate modified transformers by original Transformers: The final step is to approximate the modified transformers by the original transformer which uses ReLU and softmax.

We provide the full details in App. A.

### 3.3 Turing Completeness

Transformers are a very general class. In the original paper of Vaswani et al. [92], they were used in both an encoder and a decoder. While the previous section outlined how powerful just the encoders were, another natural question is to ask what the additional power of both a decoder along with an encoder is? Pérez et al. [73] showed that the full transformer based on a quadratic attention mechanism is Turing Complete. This result makes one unrealistic assumption, which is that the model works on arbitrary precision model. Of course, this is necessary as otherwise, Transformers are bounded finite state machines and cannot be Turing Complete.

It is natural to ask if the full attention mechanism is necessary. Or can a sparse attention mechanism also be used to simulate any Turing Machine? We show that this is indeed the case: we can use a sparse encoder and sparse decoder to simulate any Turing Machine.

To use the sparse attention mechanism in the transformer architecture, we need to define a suitable modification where each token only reacts to previous tokens. Unlike the case for BERT, where the entire attention mechanism is applied once, in full transformers, the sparse attention mechanism at decoder side is used token by token. Secondly the work of Pérez et al. [73], uses each token as a representation of the tape history and uses the full attention to move and retrieve the correct tape symbol. Most of the construction of Pérez et al. [73] goes through for sparse attentions, except for their addressing scheme to point back in history (Lemma B.4 in [73]). We show how to simulate this using a sparse attention mechanism and defer the details to App. B.

## 3.4 Limitations

We demonstrate a natural task which can be solved by the full attention mechanism in O(1)-layers. However, under standard complexity theoretic assumptions, this problem requires  $\tilde{\Omega}(n)$ -layers for any sparse attention layers with  $\tilde{O}(n)$  edges (not just BIGBIRD). (Here  $\tilde{O}$  hides poly-logarthmic factors.)

Consider the simple problem of finding the corresponding furthest vector for each vector in the given sequence of length n. Formally,

**Task 1.** Given n unit vectors  $\{u_1, \ldots, u_n\}$ , find  $f(u_1, \ldots, u_n) \to (u_{1^*}, \ldots, u_{n^*})$  where for a fixed  $j \in [n]$ , we define  $j^* = \arg \max_k \|u_k - u_j\|_2^2$ .

Finding vectors that are furthest apart boils down to minimize inner product search in case of unit vectors. For a full-attention mechanism with appropriate query and keys, this task is very easy as we can evaluate all pair-wise inner products.

The impossibility for sparse-attention follows from hardness results stemming from Orthogonal Vector Conjecture(OVC) [1, 2, 7, 97]. The OVC is a widely used assumption in fine-grained complexity. Informally, it states that one cannot determine if the minimum inner product among n boolean vectors is 0 in subquadratic time. In App. C, we show a reduction using OVC to show that if a transformer  $g \in \mathcal{T}_D^{H=1,m=2d,q=0}$  for any sparse directed graph D can evaluate the Task 1, it can solve the orthogonal vector problem.

**Proposition 1.** There exists a single layer full self-attention  $g \in \mathcal{T}^{H=1,m=2d,q=0}$  that can evaluate Task 1, i.e.  $g(u_1,...,u_n) = [u_{1^*},...,u_{n^*}]$ , but for any sparse-attention graph D with  $\tilde{O}(n)$  edges (i.e. inner product evaluations), would require  $\tilde{\Omega}(n^{1-o(1)})$  layers.

We give a formal proof of this fact in App. C.

# 4. Experiments: Natural Language Processing

In this section our goal is to showcase benefits of modeling longer input sequence for NLP tasks, for which we select three representative tasks. We begin with basic masked language modeling (MLM; Devlin et al. 23) to check if better contextual representations can be learnt by utilizing longer contiguous sequences. Next, we consider QA with supporting evidence, for which capability to handle longer sequence would allow us to retrieve more evidence using crude systems like TF-IDF/BM25. Next, we tackle long document classification where

Dataset	# tokens	Avg. doc len.
Books [110]	1.0B	37K
CC-News $[35]$	7.4B	561
Stories [90]	7.7B	8.2K
Wikipedia	3.1B	592

Model	Base	Large
RoBERTa (sqln: 512)	1.846	1.496
Longformer (sqln: 4096)	1.705	1.358
BigBird-itc (sqln: 4096)	1.678	1.456
BigBird-etc (sqln: 4096)	1.611	1.274

Table 2: Dataset used for pre training.

Table 3: MLM BPC on held-out set.

discriminating information may not be located in first 512 tokens. Finally, we present BIGBIRD for generating abstractive summaries for documents longer than 512 token, which is a very realistic scenario. Below we summarize the results for BIGBIRD using sequence length 4096, while we defer all other setup details including computational resources, batch size, step size, number of steps, etc. in App. E.

Pretraining and MLM We follow [23, 64] to create base and large versions of BIGBIRD and pretrain it using MLM objective. This task involves predicting a random subset of tokens which have been masked out. We use four standard data-sets for pretraining (listed in Tab. 2), warm-starting from the public RoBERTa checkpoint<sup>1</sup>. We compare performance in predicting the masked out tokens in terms of bits per character, following [8]. As seen in Tab. 3, both BIGBIRD and Longformer perform better than limited length RoBERTa, with BIGBIRD-ETC performing the best. We remark that Longformer was trained with a batch size of 16-32 with 48GB memory/chip, where as we trained on a more reasonable 16GB memory/chip with batch size of 32-64. Our increased memory efficiency is due to efficient blocking and sparsity structure of the sparse attention mechanism described in Sec. 2.

# 4.1 Encoder Only Task

For these tasks we only need to train encoder and we employ sparse attention mechanism of BIGBIRD as described in Sec. 2.

Question Answering (QA) We considered following four challenging datasets:

- 1. Natural Questions [53]: For the given question, find a short span of answer (SA) from the given evidences as well highlight the paragraph from the given evidences containing information about the correct answer (LA).
- 2. HotpotQA-distractor [101]: Similar to natural questions, it requires finding the answer (Ans) as well as the supporting facts (Sup) over different documents needed for multi-hop reasoning from the given evidences.
- 3. TriviaQA-wiki [42]: We need to provide an answer for the given question using provided Wikipedia evidence, however, the answer might not be present in the given evidence. On a smaller *verified* subset of question, the given evidence is guaranteed to contain the answer. Nevertheless, we model the answer as span selection problem in this case as well.
- 4. WikiHop [96]: Chose correct option from multiple-choice questions (MCQ), by aggregating information spread across multiple documents given in the evidences.

<sup>1.</sup> https://github.com/pytorch/fairseq/tree/master/examples/roberta

Model	HotpotQA			Natu	ıralQ	TriviaQA	WikiHop
1110401	Ans	Sup	Joint	LA	SA	Full	$\overline{\text{MCQ}}$
RoBERTa	73.5	83.4	63.5	-	-	74.3	72.4
Longformer	74.3	84.4	64.4	-	-	75.2	75.0
BIGBIRD-ITC	75.7	86.8	67.7	70.8	53.3	79.5	<b>75.9</b>
BigBird-etc	75.5	87.1	67.8	73.9	54.9	78.7	<b>75.9</b>

Table 4: QA Dev results using Base size models. We report accuracy for WikiHop and F1 for HotpotQA, Natural Questions, and TriviaQA.

Model	HotpotQA			Natu	NaturalQ		TriviaQA	
Woder	Ans	Ans Sup Joint LA SA		SA	Full	Verified	$\overline{\text{MCQ}}$	
HGN [27]	82.2	88.5	74.2	-	-	-	-	-
GSAN	81.6	88.7	73.9	-	-	-	-	-
ReflectionNet [33]	-	-	-	77.1	64.1	-	-	-
RikiNet [62]	-	-	-	75.5	59.5	-	-	-
Fusion-in-Decoder [40]	-	-	-	-	-	84.5	90.3	-
SpanBERT [43]	-	-	-	-	-	79.1	86.6	-
MRC-GCN [88]	-	-	-	-	-	-	-	78.3
MultiHop [14]	-	-	-	-	-	-	-	76.5
Longformer [8]	81.2	85.8	73.2	-	-	77.3	85.3	81.9
BIGBIRD-ETC	81.2	89.1	73.6	77.7	57.8	80.9	90.8	82.3

Table 5: Fine-tuning results on **Test** set for QA tasks. The Test results (F1 for HotpotQA, Natural Questions, TriviaQA, and Accuracy for WikiHop) have been picked from their respective leaderboard. For each task the top-3 leaders were picked not including BIGBIRD-etc. For Natural Questions Long Answer (LA), TriviaQA Verified, and WikiHop, BIGBIRD-ETC is the new state-of-the-art. On HotpotQA we are third in the leaderboard by F1 and second by Exact Match (EM).

As these tasks are very competitive, multiple highly engineered systems have been designed specific each dataset confirming to respective output formats. For a fair comparison, we had to use some additional regularization for training BIGBIRD, details of which are provided in App. E along with exact architecture description. We experiment using the base sized model and select the best configuration on the development set for each dataset (as reported in Tab. 4). We can see that BIGBIRD-ETC, with expanded global tokens consistently outperforms all other models. Thus, we chose this configuration to train a large sized model to be used for evaluation on the hidden test set.

In Tab. 5, we compare BIGBIRD-ETC model to top-3 entries from the leaderboard excluding BIGBIRD. One can clearly see the importance of using longer context as both Longformer and BIGBIRD outperform models with smaller contexts. It is worth noting that BIGBIRD submission is a single model, whereas the other top-3 entries for Natural Questions are ensembles, which might explain the slightly lower accuracy in exact answer phrase selection. Also, high performing Fusion-in-Decoder [40] model for TriviaQA can be considered to be

Model	IMDb [65]	Yelp-5 [108]	Arxiv [36]	Patents [54]	Hyperpartisan [48]
# Examples	25000	650000	30043	1890093	645
# Classes	2	5	11	663	2
Excess fraction	0.14	0.04	1.00	0.90	0.53
SoTA	[89] 97.4	[3] 73.28	[70] 87.96	[70] 69.01	[41] 90.6
RoBERTa	$95.0 \pm 0.2$	71.75	87.42	67.07	$87.8 \pm 0.8$
BigBird	$95.2 \pm 0.2$	72.16	92.31	69.30	$92.2 \pm 1.7$

Table 6: Classification results. We report the F1 micro-averaged score for all datasets. Experiments on smaller IMDb and Hyperpartisan datasets are repeated 5 times and the average performance is presented along with standard deviation.

a sparse long sequence encoder-decoder transformer, despite being motivated differently in their paper. In particular, their sparse attention at encoder side would be block diagonal, thereby being a special case of BigBird with r = 0, g = 0.

Document Classification We experiment on datasets of different lengths and contents as listed in tab. 6. We compute the fraction of the dataset that exceeds 512 tokens, i.e. the length at which the document are often truncated. Following BERT, we used one layer with cross entropy loss on top of the first [CLS] token. We see that gains of using BIGBIRD are more significant when we have longer documents and fewer training examples. For instance, using base sized model, BIGBIRD improves state-of-the-art for Arxiv dataset by about 5% points. On Patents dataset, there is improvement over using simple BERT/RoBERTa, but given the large size of training data the improvement over SoTA is not significant. Note that this performance gain is not seen for much smaller IMDb dataset. Along with experimental setup detail, we present results on GLUE tasks in app. E which show competitive performance.

#### 4.2 Encoder-Decoder Tasks

For an encoder-decoder setup, one can easily see that both suffer from quadratic complexity due to the full self attention. We focus on introducing the sparse attention mechanism of BIGBIRD only at the encoder side. This is because, in practical generative applications, the length of output sequence is typically small as compared to the input. For example for text summarization, we see in Tab. 7 that the median output sequence length is  $\sim 200$  where as the input sequence's median length is > 3000. For such applications, it is more efficient to use sparse attention mechanism for the encoder and full self-attention for the decoder.

	Instances				Input Length			Output Length		
Dataset	Training	Dev	Test	_	Median	90%-ile	-	Median	90%-ile	
Arxiv [21]	203037	6436	6440		6151	14405		171	352	
PubMed [21]	119924	6633	6658		2715	6101		212	318	
BigPatent [79]	1207222	67068	67072		3082	7693		123	197	

Table 7: Statistics of datasets used for summarization.

	Model		Arxiv		$\operatorname{PubMed}$			]	BigPatent		
Μ			R-2	R-L	R-1	R-2	R-L	R-1	R-2	R-L	
	SumBasic [69]	29.47	6.95	26.30	37.15	11.36	33.43	27.44	7.08	23.66	
	LexRank [26]	33.85	10.73	28.99	39.19	13.89	34.59	35.57	10.47	29.03	
	LSA [98]	29.91	7.42	25.67	33.89	9.93	29.70	-	-	-	
r	Attn-Seq2Seq [86]	29.30	6.00	25.56	31.55	8.52	27.38	28.74	7.87	24.66	
Art	Pntr-Gen-Seq2Seq [78]	32.06	9.04	25.16	35.86	10.22	29.69	33.14	11.63	28.55	
rior	Long-Doc-Seq2Seq [21]	35.80	11.05	31.80	38.93	15.37	35.21	-	-	-	
$P_{ m I}$	Sent-CLF [82]	34.01	8.71	30.41	45.01	19.91	41.16	36.20	10.99	31.83	
	Sent-PTR [82]	42.32	15.63	38.06	43.30	17.92	39.47	34.21	10.78	30.07	
	Extr-Abst-TLM [82]	41.62	14.69	38.03	42.13	16.27	39.21	38.65	12.31	34.09	
	Dancer [32]	42.70	16.54	38.44	44.09	17.69	40.27	-	-	-	
	Transformer	28.52	6.70	25.58	31.71	8.32	29.42	39.66	20.94	31.20	
se	+ RoBERTa [77]	31.98	8.13	29.53	35.77	13.85	33.32	41.11	22.10	32.58	
Base	+ Pegasus [107]	34.81	10.16	30.14	39.98	15.15	35.89	43.55	20.43	31.80	
	BIGBIRD-RoBERTa	41.22	16.43	36.96	43.70	19.32	39.99	55.69	37.27	45.56	
.e	Pegasus (Reported) [107]	44.21	16.95	38.83	45.97	20.15	41.34	52.29	33.08	41.75	
arge	Pegasus (Re-eval)	43.85	16.83	39.17	44.53	19.30	40.70	52.25	33.04	41.80	
Ļ	BigBird-Pegasus	46.63	19.02	41.77	46.32	20.65	42.33	60.64	42.46	50.01	

Table 8: Summarization ROUGE score for long documents.

Summarization Document summarization is a task of creating a short and accurate summary of a text document. We used three long document datasets for testing our model details of which are mention in Tab. 7. In this paper we focus on abstractive summarization of long documents where using a longer contextual encoder should improve performance. The reasons are two fold: First, the salient content can be evenly distributed in the long document, not just in first 512 tokens, and this is by design in the BigPatents dataset [79]. Second, longer documents exhibit a richer discourse structure and summaries are considerably more abstractive, thereby observing more context helps. As has been pointed out recently [77, 107], pretraining helps in generative tasks, we warm start from our general purpose MLM pretraining on base-sized models as well as utilizing state-of-the-art summarization specific pretraining from Pegasus [107] on large-sized models. The results of training BigBird sparse encoder along with full decoder on these long document datasets are presented in Tab. 8. We can clearly see modeling longer context brings significant improvement. Along with hyperparameters, we also present results on shorter but more widespread datasets in App. E.4, which show that using sparse attention does not hamper performance either.

## 5. Experiments: Genomics

There has been a recent upsurge in using deep learning for genomics data [13, 87, 106], which has resulted in improved performance on several biologically-significant tasks such as promoter site prediction [72], methylation analysis [56], predicting functional effects of non-coding variant [109], etc. These approaches consume DNA sequence fragments as inputs,

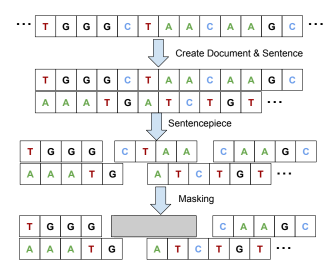


Figure 2: Visual description of how the masked language modeling data was generated from raw DNA dataset. The raw DNA sequences of GRCh37, where split at random positions to create documents with 50-100 sentences where each sentence was 500-1000 base pairs (bps). Thus each document had a continuous strand of 25000-100,000 bps of DNA. This process was repeated 10 times to create 10 sets of document for each chromosome of GRCH37. The resulting set of documents was then passed through Sentencepiece that created tokens of average 8bp. For pretraining we used masked language model and masked 10% of the tokens and trained on predicting the masked tokens.

and therefore we believe longer input sequence handling capability of BIGBIRD would be beneficial as many functional effects in DNA are highly non-local [12]. Furthermore, taking inspiration from NLP, we learn powerful contextual representations for DNA fragments utilizing abundant unlabeled data (e.g. human reference genome, Saccharomyces Genome Database) via MLM pretraining. Next, we showcase that our long input BIGBIRD along with the proposed pretraining significantly improves performances in two downstream tasks. Detailed experimental setup for the two tasks are provided in App. F.

Pre-training and MLM Instead of operating directly on base pairs, we propose to first segment DNA into tokens so as to further increase the context length (Fig. 2). This idea is not entirely novel and has been previously explored in Liang [59]. In particular, we build a byte-pair encoding [51] table for the DNA sequence of size 32K, with each token representing 8.78 base pairs on average. We learn contextual representation of

Model	BPC
SRILM [59] BERT (sqln. 512)	1.57 1.23
BIGBIRD (sqln. 4096)	1.12

Table 9: MLM BPC

these token on the human reference genome (GRCh37)<sup>2</sup> using MLM objective.

We then report the bits per character (BPC) on a held-out set in Tab. 9. We find that attention based contextual representation of DNA does improve BPC, which is further improved by using longer context in BIGBIRD.

<sup>2.</sup> https://www.ncbi.nlm.nih.gov/assembly/GCF\_000001405.13/

Promoter Region Prediction Promoter is a DNA region typically located upstream of the gene, which is the site of transcription initiation. Multiple methods have been proposed to identify the promoter regions in a given DNA sequence [100, 60, 11, 99, 72], as it is an important first step in understanding gene regulation. The corresponding machine learning task is to classify a given DNA fragment as promoter or non-promoter sequence. We use

Model	F1
CNNProm [91] DeePromoter [72]	69.7 95.6
BIGBIRD	99.9

Table 10: Comparison.

the dataset compiled by Oubounyt et al. [72] which was built from Eukaryotic Promoter Database (EPDnew) [25] <sup>3</sup>. We finetuned the pretrained BIGBIRD model from above, using the training data and report F1 on test dataset. We compare our results to the previously reported best method in Tab. 10. We see that BIGBIRD achieve nearly perfect accuracy with a 5% jump from the previous best reported accuracy.

Chromatin-Profile Prediction Non-coding regions of DNA do not code for proteins. Majority of diseases and other trait associated single-nucleotide polymorphism are correlated to non-coding genomic variations [109, 47]. Thus, understanding the functional effects of non-coding regions of DNA is a very important task. An important step in this process, as defined by Zhou and Troyanskaya [109], is to predict large-scale chromatin-

Model	TF	HM	DHS
gkm-SVM [31] DeepSea [109]	89.6 95.8	85.6	92.3
BIGBIRD	96.1	88.7	92.1

Table 11: Chromatin-Profile Prediction

profiling from non-coding genomic sequence. To this effect, DeepSea [109], compiled 919 chromatin-profile of 2.4M non-coding variants from Encyclopedia of DNA Elements (ENCODE)<sup>4</sup> and Roadmap Epigenomics projects<sup>5</sup>. The corresponding ML task is to predict, for a given non-coding region of DNA, these 919 chromatin-profile including 690 transcription factors (TF) binding profiles for 160 different TFs, 125 DNase I sensitivity (DHS) profiles and 104 histone-mark (HM) profiles. We jointly learn 919 binary classifiers to predict these functional effects from sequence of DNA fragments. On held-out chromosomes, we compare AUC with the baselines in Tab. 11 and see that we significantly improve on performance on the harder task HM, which is known to have longer-range correlations [28] than others.

### 6. Conclusion

We propose BigBird: a sparse attention mechanism that is linear in the number of tokens. BigBird satisfies a number of theoretical results: it is a universal approximator of sequence to sequence functions and is also Turing complete. Theoretically, we use the power of extra global tokens preserve the expressive powers of the model. We complement these results by showing that moving to sparse attention mechanism do incur a cost. Empirically, BigBird gives state-of-the-art performance on a number of NLP tasks such as question answering and long document summarization. We further introduce attention based contextual language model for DNA and fine-tune it for down stream tasks such as promoter region prediction and predicting effects of non-coding variants.

<sup>3.</sup> https://epd.epfl.ch/human/human\_database.php?db=human

<sup>4.</sup> https://www.encodeproject.org/

<sup>5.</sup> http://www.roadmapepigenomics.org/

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# Big Bird: Bert for Longer Sequences: Appendix

# Appendix A. Universal Approximators

#### A.1 Notation

We begin by setting up some notations following Pérez et al. [73] to formally describe the complete architecture of Transformers. A single layer of Transformer encoder is a parametric function Enc receiving a sequence  $\boldsymbol{X} = (\boldsymbol{x}_1, ..., \boldsymbol{x}_n)$  of vectors in  $\mathbb{R}^d$  and returning a sequence  $\boldsymbol{Z} = (\boldsymbol{z}_1, ..., \boldsymbol{z}_n)$  of the same length. Each  $\boldsymbol{z}_i$  is a d dimensional vector as well. We interchangeably treat the sequence  $\boldsymbol{X}$  as a matrix in  $\mathbb{R}^{n \times d}$ . Enc has two components:

- 1. An attention mechanism ATTN that takes in the sequence X and returns sequence  $(a_1, ..., a_n)$  of the same length and dimensionality; and
- 2. A two layer fully connected network O that takes in a vector in  $\mathbb{R}^d$  and returns a vector in  $\mathbb{R}^d$ .

Then *i*-th output vector of Enc(X) is computed as follows:

$$z_i = O(a_i) + a_i$$
 where  $a_i = ATTN(X)_i + x_i$  (1)

Now it remains to define ATTN and O which we do next.

As described in Sec. 2, an attention mechanism is parameterized by three functions: Q, K, V:  $\mathbb{R}^d \to \mathbb{R}^m$ . In this paper, we assume that they are simply matrix products:  $Q(\boldsymbol{x}) = \boldsymbol{x}W_Q$ ,  $K(\boldsymbol{x}) = \boldsymbol{x}W_K$ , and  $V(\boldsymbol{x}) = \boldsymbol{x}W_V$ , where  $W_Q, W_K, W_V \in \mathbb{R}^{d \times m}$  and  $W_V \in \mathbb{R}^{d \times d}$ . In reality a multi-headed attention is used, i.e. we have not only one, but H-sets of Query/Key/Value weight matrices,  $W_Q^h, W_V^h, W_K^h$  for h = 1, ..., H. Thus, for a directed graph D over [n], the i<sup>th</sup> output vector of the generalized attention mechanism would be

$$ATTN_D(\boldsymbol{X})_i = \sum_{h=1}^{H} \sigma\left((\boldsymbol{x}_i W_Q^h) (\boldsymbol{X}_{N(i)} W_K^h)^T\right) \cdot (\boldsymbol{X}_{N(i)} W_V^h)$$
(AT)

where N(i) denote the out-neighbors set of node i in D. In other words, the set of arcs (directed edges) in D represents the set of inner products that our attention mechanism will consider. Also recall that  $\sigma$  is a scoring function such as softmax or hardmax.

Lastly, we define the output fully connected network as follows:

$$O(\mathbf{a}_i) = \text{ReLU}(\mathbf{a}_i W_1 + b_1) W_2 \cdot + b_2$$
 (FF)

Here  $W_1 \in \mathbb{R}^{d \times q}$ ,  $W_2 \in \mathbb{R}^{q \times d}$ ,  $b_1 \in \mathbb{R}^p$ , and  $b_2 \in \mathbb{R}^d$  are parameters of output network O.

Additional Notation We introduce a few pieces of additional notation that will be useful. Let  $[a,b)_{\delta} = \{a,a+\delta,\ldots,a+\lfloor\frac{b-a}{\delta}\rfloor\cdot\delta\}$ . Therefore,  $[0,1)_{\delta} = \{0,\delta,2\delta,\ldots,(1-\delta)\}$ . We use  $\mathbf{1}[\mathcal{E}]$  to denote the indicator variable; it is 1 if the event  $\mathcal{E}$  occurs and 0 otherwise.

# A.2 Proof

In this section, we will present the full proof of theorem 1. The proof will contain three parts. The first and the third part will largely follow standard techniques. The main innovation lies is in the second part.

# A.2.1 Approximate $\mathcal{F}_{CD}$ by Piece-wise constant functions

First, we consider a suitable partition of the region (0,1) into a grid of granularity  $\delta$ , which we denote by  $G_{\delta}$ . We do this using Lemma 8 from Yun et al. [105], which we restate for completeness:

**Lemma 1** (Lemma 8 [105]). For any given  $f \in \mathcal{F}_{CD}$  and  $1 \leq p \leq \infty$ , there exists a  $\delta > 0$  such that there exists a piece-wise constant function  $\bar{f}$  with  $d_p(f,\bar{f}) \leq \frac{\epsilon}{3}$ . Concretely,  $\bar{f}$  is defined as

$$\bar{f}(X) = \sum_{P \in \mathbb{G}_{\delta}} f(P) \cdot \mathbf{1} \left[ \| \text{ReLU}(X - P) \|_{\infty} \le \delta \right]$$

Since transformers can learn a positional embedding E, without any loss of generality, we can consider the translated function. In particular, define

$$E = \begin{bmatrix} 0 & 0 & 0 & \dots & 0 \\ \delta^{-d} & \delta^{-d} & \delta^{-d} & \dots & \delta^{-d} \\ \delta^{-2d} & \delta^{-2d} & \delta^{-2d} & \dots & \delta^{-2d} \\ \vdots & & & & & \\ \delta^{-(n-1)d} & \delta^{-(n-1)d} & \delta^{-(n-1)d} & \dots & \delta^{-(n-1)d} \end{bmatrix}$$

We will try to approximate g(X) = f(X - E) where g is defined on the domain  $[0,1]^d \times [\delta^{-d}, \delta^{-d} + 1]^d \times \cdots \times [\delta^{-(n-1)d}, \delta^{-(n-1)d} + 1]^d$ . To do so, we will apply a suitable modification of Lemma 1, which will consider the discretized grid

$$\mathbf{G}_{\delta}^{E} := [0, 1]_{\delta}^{d} \times [\delta^{-d}, \delta^{-d} + 1]_{\delta}^{d} \times \dots \times [\delta^{-(n-1)d}, \delta^{-(n-1)d} + 1]_{\delta}^{d}.$$

Therefore, it suffices to approximate a function  $\bar{f}: \mathbf{G}_{\delta}^{E} \to \mathbb{R}^{n \times d}$  defined as

$$\bar{f}(X) = \sum_{P \in \mathbf{G}_{\delta}^{E}} f(P - E) \cdot \mathbf{1} \left[ \| \operatorname{ReLU}(X - P) \|_{\infty} \le \delta \right].$$

# A.2.2 CONTEXTUAL MAPPINGS AND SPARSE ATTENTION MECHANISMS

Throughout this section, we will assume that we are given a function that has an extra global token at index 0 and all vectors have an extra dimension appended to them. The latter assumption is without loss of generality as we can use the Feed-Forward Network to append sparse dimensions. In particular, we will associate  $X \in \mathbb{R}^{(n+1)\times(d+1)}$  where we write  $X = (x_0, x_1, \ldots, x_n)$ . Although our function is only defined for  $\mathbf{G}_{\delta}^E \subset \mathbb{R}^{n \times d}$ , we can amend

the function in a natural way by making it ignore the first column. To avoid excessive clutter, we will assume that the function value is evaluated on the last n columns.

The main idea in this section is the use of contextual mapping to enable Transformers to compute any discretized function. A contextual mapping is an unique encoding of each tuple  $(X, x_i)$  where  $X \in \mathbf{G}_{\delta}^E$ , and each column  $x_i \in [\delta^{-(i-1)d}, \delta^{-(i-1)d} + 1)_{\delta}^d$  for all  $i \in [n]$ . We restate the definition adapted to our setting below

**Definition 2** (Defn 3.1 [105]). (Contextual Mapping) A contextual mapping is a function mapping  $q: \mathbf{G}_{\delta}^{E} \to \mathbb{R}^{n}$  if it satisfies the following:

- 1. For any  $P \in \mathbf{G}_{\delta}^{E}$ , q(P) contains distinct entries.
- 2. For any two  $P, P' \in \mathbf{G}_{\delta}^{E}$  with  $P \neq P'$ , all entries of q(P) and q(P') are distinct.

The key technical novelty of the proof is computing a contextual mapping using only the sparse attention mechanism. We create a "selective shift" operator which only shifts entries of a vector that lie in a certain range. We will use this shift operator strategically to ensure that we attain a contextual mapping at the end of the process. The lemma below, which is based on parts of the proof of Lemma 6 of [105], states that we can implement a suitable "selective" shift operator using a sparse attention mechanism.

**Lemma 2.** Given a function  $\psi : \mathbb{R}^{(n+1)\times(d+1)} \times \mathbb{R}^2 \to \mathbb{R}^{(n+1)\times 1}$  and a vector  $u \in \mathbb{R}^{d+1}$  and a sparse attention mechanism based on the directed graph D, we can implement a selective shift operator that receives as input a matrix  $X \in \mathbb{R}^{(n+1)\times(d+1)}$  and outputs  $X + \rho \cdot \psi_u(X, b_1, b_2)$  where

$$\psi_u(Z; b_1, b_2)_i = \begin{cases} (\max_{j \in N(i)} u^T Z_j - \min_{j \in N(i)} u^T Z_j) e_1 & \text{if } b_1 \le u^T Z_j \le b_2 \\ 0 & \text{else.} \end{cases}$$

Note that  $e_1 \in \mathbb{R}^{d+1}$  denotes  $(1, 0, \dots, 0)$ .

*Proof.* Consider the function, which can be implemented by a sparse attention mechanism:

$$\tilde{\psi}(X,b)_i = \sigma_H \left[ (u^T \cdot X_i)^T \cdot (u^T X_{N(i)} - b 1_{N(i)}^T) e^{(1)} (u^T X_{N(i)}) \right]$$

This is because the Key, Query and Value functions are simply affine transformations of X. Given any graph D, the above function will evaluate to the following:

$$\tilde{\psi}(Z;b)_i = \begin{cases} (\max_{j \in N(i)} u^T Z_j) e_1 & \text{if } u^T Z_j > b \\ (\min_{j \in N(i)} u^T Z_j) e_1 & \text{if } u^T Z_j < b \end{cases}$$

Therefore we can say that  $\tilde{\psi}(Z; b_Q) - \tilde{\psi}(Z; b_{Q'})$  satisfies

$$\psi(Z; b_1, b_2)_i = \begin{cases} (\max_{j \in N(i)} u^T Z_j - \min_{j \in N(i)} u^T Z_j) e_1 & \text{if } b_1 \le u^T Z_j \le b_2 \\ 0 & \text{else} \end{cases}$$

The following lemma, which is the heart of the proof, uses the above selective shift operators to construct contextual mappings.

**Lemma 3.** There exists a function  $g_c : \mathbb{R}^{(n+1)\times(d+1)} \to \mathbb{R}^{(n+1)}$  and a unique vector u, such that for all  $P \in \mathbf{G}_{\delta}^E$   $g_c(P) := \langle u, g(P) \rangle$  satisfies the property that  $g_c$  is a contextual mapping of P. Furthermore,  $g_c \in \mathcal{T}_D^{2,1,1}$  using a composition of sparse attention layers as long as D contains the star graph.

*Proof.* Define  $u \in \mathbb{R}^{d+1} = [1, \delta^{-1}, \delta^{-2}, \dots, \delta^{-d+1}, \delta^{-nd}]$  and let  $X_0 = (0, \dots, 0, 1)$ . We will assume that  $\langle x_i, x_0 \rangle = 0$ , by assuming that all the columns  $x_1, \dots, x_n$  are appended by 0.

To successfully encode the entire context in each token, we will interleave the shift operator to target the original columns  $1, \ldots, n$  and to target the global column 0. After a column i is targeted, its inner product with u will encode the entire context of the first i columns. Next, we will shift the global token to take this context into account. This can be subsequently used by the remaining columns.

For  $i \in \{0, 1, ..., n\}$ , we will use  $l_i$  to denote the innerproducts  $\langle u, x_i \rangle$  at the beginning. For  $f_i = \langle u, x_i \rangle$  after the  $i^{th}$  column has changed for  $i \in \{1, ..., n\}$  and we will use  $f_0^k$  to denote  $\langle u, x_0 \rangle$  after the  $k^{th}$  phase. We need to distinguish the global token further as it's inner product will change in each phase. Initially, given  $X \in \mathbf{G}_{\delta}^E$ , the following are true:

$$\delta^{-(i-1)d} \le \langle u, X_i \rangle \le \delta^{-id} - \delta \qquad \text{for all } i \in [n]$$
$$\delta^{-(n+1)d} = \langle u, X_0 \rangle$$

Note that all  $l_i$  ordered in distinct buckets  $l_1 < l_2 < \cdots < l_n < l_0$ .

We do this in phases indexed from  $i \in \{1, ..., n\}$ . Each phase consists of two distinct parts: **The low shift operation:** These operation will be of the form

$$X \leftarrow X + \delta^{-d}\psi\left(X, v - \delta/2, v + \delta/2\right)$$

for values  $v \in [\delta^{-id}), \delta^{-(i+1)d})_{\delta}$ . The range is chosen so that only  $l_i$  will be in the range and no other  $l_j$   $j \neq i$  is in the range. This will shift exactly the  $i^{th}$  column  $x_i$  so that the new inner product  $f_i = \langle u, x_i \rangle$  is substantially larger than  $l_i$ . Furthermore, no other column of X will be affected.

The high shift operation: These operation will be of the form

$$X \leftarrow X + \delta^{-nd} \cdot \psi (X, v - \delta/2, v + \delta/2)$$

for values  $v \in [S_i, T_i)_{\delta}$ . The range  $[S_i, T_i)_{\delta}$  is chosen to only affect the column  $x_0$  (corresponding to the global token) and no other column. In particular, this will shift the global token by a further  $\delta^{-nd}$ . Let  $\tilde{f}_0^i$  denote the value of  $\tilde{f}_0^i = \langle u, x_0 \rangle$  at the end of  $i^{th}$  high operation.

Each phase interleaves a shift operation to column i and updates the global token. After each phase, the updated  $i^{th}$  column  $f_i = \langle u, x_i \rangle$  will contain a unique token encoding the values of all the  $l_1, \ldots, l_i$ . After the high update,  $\tilde{f}_0^i = \langle u, x_0 \rangle$  will contain information about the first i tokens.

Finally, we define the following constants for all  $k \in \{0, 1, ..., n\}$ .

$$T_k = (\delta^{-(n+1)d} + 1)^k \cdot \delta^{-nd} - \sum_{t=2}^k (\delta^{-(n+1)d} + 1)^{k-t} (2\delta^{-nd-d} + \delta^{-nd} + 1)\delta^{-td}$$
$$- (\delta^{-(n+1)d} + 1)^{k-1} (\delta^{-nd-d} + \delta^{-nd})\delta^{-d} - \delta^{-(k+1)d}$$
(UP)

$$S_k = (\delta^{-(n+1)d} + 1)^k \cdot \delta^{-nd} - \sum_{t=2}^k (\delta^{-(n+1)d} + 1)^{k-t} (2\delta^{-nd-d} + \delta^{-nd} + 1)\delta^{-(t-1)d} - (\delta^{-(n+1)d} + 1)^{k-1} (\delta^{-nd-d} + \delta^{-nd}) - \delta^{-kd}$$
(LP)

After each k phases, we will maintain the following invariants:

- 1.  $S_k < \tilde{f}_0^k < T_k \text{ for all } k \in \{0, 1, \dots, n\}.$
- 2.  $T_{k-1} < f_k < S_k$
- 3. The order of the inner products after  $k^{th}$  phase is

$$l_{k+1} < l_{k+2} \cdots < l_n < f_1 < f_2 < \cdots < f_k < \tilde{f}_0^k$$

Base case The case k = 0, is trivial as we simply set  $S_0 = \delta^{-(n+1)d}$ ,  $T_0 = \delta^{-(n+1)d} + \delta$ . The first nontrivial case is k = 1.

Inductive Step First, in the low shift operation is performed in the range  $[\delta^{-(k-1)d}, \delta^{-kd})_{\delta}$  Due to the invariant, we know that there exists only one column  $x_k$  that is affected by this shift. In particular, for column k, we will have  $\max_{j \in N(k)} \langle u, x_j \rangle = \langle u, x_0 \rangle = \tilde{f}_0^{k-1}$ . The minimum is  $l_k$ . Thus the update will be  $f_k = \delta^{-d}(\tilde{f}_0^{k-1} - l_k) + l_k$ . Observe that for small enough  $\delta$ ,  $f_k \geq \tilde{f}_0^{k-1}$ . Hence the total ordering, after this operation is

$$l_k + 1 < l_{k+2} \dots < l_n < f_1 < f_2 < \dots < \tilde{f}_0^{k-1} < f_k$$
 (2)

Now when we operate a higher selective shift operator in the range  $[S_{k-1}, T_{k-1})_{\delta}$ . Since only global token's innerproduct  $\tilde{f}_0^{k-1}$  is in this range, it will be the only column affected by the shift operator. The global token operates over the entire range, we know from Eq. (2) that,  $f_k = \max_{i \in [n]} \langle u, x_i \rangle$  and  $l_{k+1} = \min_{i \in [n]} \langle u, x_i \rangle$ . The new value  $\tilde{f}_0^k = \delta^{-nd} \cdot (f_k - l_{k+1}) + \tilde{f}_0^{k-1}$ . Expanding and simplifying we get,

$$\begin{split} \tilde{f}_0^k &= \delta^{-nd} \cdot (f_k - l_{k+1}) + \tilde{f}_0^{k-1} \\ &= \delta^{-nd} \cdot (\delta^{-d} (\tilde{f}_0^{k-1} - l_k) + l_k - l_{k+1}) + \tilde{f}_0^{k-1} \\ &= \delta^{-(n+1)d} \cdot (\tilde{f}_0^{k-1} - l_k) + \delta^{-nd} (l_k - l_{k+1}) + \tilde{f}_0^{k-1} \\ &= (\delta^{-(n+1)d} + 1) \tilde{f}_0^{k-1} - (\delta^{-nd-d} + \delta^{-nd}) l_k - l_{k+1} \end{split}$$

Expanding the above recursively, we get

$$= (\delta^{-(n+1)d} + 1)^k \cdot \tilde{f}_0^0 - \sum_{t=2}^k (\delta^{-(n+1)d} + 1)^{k-t} (2\delta^{-nd-d} + \delta^{-nd} + 1) l_t$$
$$- (\delta^{-(n+1)d} + 1)^{k-1} (\delta^{-nd-d} + \delta^{-nd}) l_1 - l_{k+1}$$

Since we know that  $\tilde{f}_0^0 = \delta^{-nd}$  and each  $l_i < \delta^{-id}$ , we can substitute this to get Eq. (UP) and we can get an lower-bound Eq. (LP) by using  $l_i \ge \delta^{-(i-1)d}$ .

By construction, we know that  $S_k \leq \tilde{f}_0^k < T_k$ . For sufficiently small  $\delta$ , observe that  $S_k \leq \tilde{f}_0^k < T_k$  all are essentially the dominant term  $\approx O(\delta^{-n(k+1)d-kd})$  and all the lower order terms do not matter. As a result it is immediate to see that that  $f_k > \delta^{-d}(\tilde{f}_0^{k-1} - l_k) > T_{k-1}$  and hence we can see that the invariant 2 is also satisfied. Since only column k and the global token are affected, we can see that invariant 3 is also satisfied.

After n iterations,  $\tilde{f}_0^n$  contains a unique encoding for any  $P \in \mathbf{G}_{\delta}^E$ . To ensure that all tokens are distinct, we will add an additional layer  $X = X + \delta^{-n^2 d} \psi(X, v - \delta/2, v + \delta/2)$  for all  $v \in [S_1, T_n)_{\delta}$ . This ensures that for all  $P, P' \in \mathbf{G}_{\delta}^E$ , each entry of q(P) and q(P') are distinct.

The previous lemma shows that we can compute a contextual mapping using only sparse transforms. We now use the following lemma to show that we can use a contextual mapping and feed-forward layers to accurately map to the desired output of the function  $\bar{f}$ .

**Lemma 4** (Lemma 7 [105]). Let  $g_c$  be the function in Lemma 3, we can construct a function  $g_v : \mathbb{R}^{(n+1)\times(d+1)} \to \mathbb{R}^{(n+1)\times d}$  composed of  $O(n\delta^{-nd})$  feed-forward layers (with hidden dimension q=1) with activations in  $\Phi$  such that  $g_v$  is defined as  $g_v(Z) = [g_v^{tkn}(Z_1), \ldots, g_v^{tkn}(Z_n)]$ , where for all  $j \in \{1, \ldots, n\}$ ,

$$g_v^{tkn}(g_c(L)_j) = f(L)_j$$

#### A.2.3 Approximating modified Transformers by Transformers

The previous section assumed we used Transformers that used hardmax operator  $\sigma_H$  and activations functions belonging to the set  $\Phi$ . This is without loss of generality as following lemma shows.

**Lemma 5** (Lemma 9 [105]). For each  $g \in \bar{\mathcal{T}}^{2,1,1}$  and  $1 \leq p \leq \infty$ ,  $\exists g \in \mathcal{T}^{2,1,4}$  such that  $d_p(g,\bar{g}) \leq \epsilon/3$ 

Combining the above lemma with the Lemma 3, we get our main result:

**Theorem 2.** Let  $1 \le p \le \infty$  and  $\epsilon > 0$ , there exists a transformer network  $g \in \mathcal{T}_D^{2,1,4}$  which achieves a ratio of  $d_p(f,g) \le \epsilon$  where D is the sparse graph.

Since the sparsity graph associated with BIGBIRD contains a star network, we know that it can express any continuous function from a compact domain.

# Appendix B. Turing Completenes

In this section, we will extend our results to the setting of Pérez et al. [73]. Our exposition will largely use their proof structure but we will make a few changes. We repeat some of the lemmas with the amendments to make the exposition self-contained.

## **B.1 Notation**

Transformer Decoder We need both an encoder and a decoder in the transformer for simulating a Turing machine. We utilize the same notation used in App. A.1 for encoders. The decoder is similar to an encoder but with additional attention to an external pair of key-value vectors ( $\mathbf{K^e} \in \mathbb{R}^{n \times m}, \mathbf{V^e} \in \mathbb{R}^{n \times d}$ ), which usually come from the encoder stack. A single layer of Transformer decoder is a parametric function Dec receiving a sequence  $\mathbf{Y}_j = (\mathbf{y}_1, \dots, \mathbf{y}_j)$  of vectors in  $\mathbb{R}^d$  plus the external ( $\mathbf{K^e}, \mathbf{V^e}$ ) and returning a sequence of vectors  $\mathbf{Z}_j = (\mathbf{z}_1, \dots, \mathbf{z}_j)$  of the same length. Each  $\mathbf{z}_i$  is a d dimensional vector as well. Dec has three components, one more than Enc:

- 1. An attention mechanism ATTN that takes in the sequence  $Y_j$  and returns sequence  $(p_1, ..., p_j)$  of the same length and dimensionality;
- 2. A cross-attention mechanism CROSSATTN that takes in the sequence  $(\boldsymbol{p}_1,...,\boldsymbol{p}_j)$  plus the external  $(\boldsymbol{K}^{\mathbf{e}},\boldsymbol{V}^{\mathbf{e}})$  and returns sequence  $(\boldsymbol{a}_1,...,\boldsymbol{a}_j)$ , with each  $\boldsymbol{a}_i \in \mathbb{R}^d$ ; and
- 3. A two layer fully connected network O that takes in a vector in  $\mathbb{R}^d$  and returns a vector in  $\mathbb{R}^d$ .

Then *i*-th output vector of  $Dec(Y_i; K^e, V^e)$  is computed as follows:

$$z_i = O(a_i) + a_i \tag{3}$$

where

$$a_i = \text{CrossAttn}(\boldsymbol{p}_i, \boldsymbol{K}^{\mathbf{e}}, \boldsymbol{V}^{\mathbf{e}}) + \boldsymbol{p}_i$$
 (4)

and

$$\boldsymbol{p}_i = \operatorname{ATTN}_D(\boldsymbol{Y}_j)_i + \boldsymbol{y}_i \tag{5}$$

Attn<sub>D</sub> and O are as defined in App. A.1 and it remains to define CrossAttn. The  $i^{\text{th}}$  output vector of multi-head cross-attention attention is given by

CROSSATTN
$$(\mathbf{Y}_j)_i = \sum_{h=1}^H \sigma\left((\mathbf{y}_i W_Q^h) (\mathbf{K}^{(e)} W_K^h)^T\right) \cdot (\mathbf{V}^{(e)} W_V^h)$$
 (6)

where  $W_Q^h, W_K^h, W_V^h \in \mathbb{R}^{d \times m}$ ,  $W_V^h \in \mathbb{R}^{d \times d}$ , for all  $h = 1, \dots H$  heads.

**Turning Machine** We will use the same setup of Turning Machine that was used by Pérez et al. [73] (see section B.4). Given a Turing Machine  $M = (Q, \Sigma, \delta, q_{init}, F)$ , we use the following notation

 $q^{(j)}$ : state of Turing machine M at time j.

 $s^{(j)}$ : symbol under the head of M at time j.

 $v^{(j)}$ : symbol written by M at time j.

 $m^{(j)}$ : head direction in the transition of M at time j.

Vector representations For a symbol  $s \in \Sigma$ ,  $[\![ s ]\!]$  denotes its one-hot vector representation in  $\mathbb{Q}^{|\Sigma|}$ . All the transformer intermediate vectors used in our simulations have dimension  $d = 2|Q| + 4|\Sigma| + 16$ . Note that we use five extra dimension as compared to Pérez et al. [73]. We follow the convention used in Pérez et al. [73] and write a a vector  $\mathbf{v} \in \mathbb{Q}^d$  arranged in four groups of values as follows

where  $\mathbf{q}_i \in \mathbb{Q}^{|Q|}$ ,  $\mathbf{s}_i \in \mathbb{Q}^{|\Sigma|}$ , and  $x_i \in \mathbb{Q}$ .

### B.2 Details of the Simulation

In this section, we give more details on the architecture of the encoder and decoder needed to implement our simulation strategy.

**High Level Overview:** Given the Turing machine M, we will show that a transformer with an appropriate encoder and decoder  $\mathcal{T}_D$  can simulate each step of M's execution.

Our simulation strategy will mostly follow Pérez et al. [73], except we will use a sparse attention mechanism. The main idea is to maintain the current Turing machine state  $q^{(j)}$  and symbol under the head  $s^{(j)}$  as part of the decoder sequence  $\mathbf{Y}$  for all time step j so that we can always simulate the corresponding Turing machine transition  $\delta(q^{(j)}, s^{(j)}) = (q^{(j)}, v^{(j)}, m^{(j)})$ . The key difference will rise in Lemma B.4 of Pérez et al. [73], where full attention is used to select the appropriate symbol from tape history in one step. To accomplish the same task with sparse attention, we will exploit the associative property of max and break down the symbol selection over multiple steps. Thus, unlike Pérez et al. [73] one decoding step of our sparse transformer  $\mathcal{T}_D$  does not correspond to one step of the Turing machine M. In particular, we will have two type of steps: compute step corresponding to update of M's state and intermediate steps corresponding to aggregating the max (which in turn is used for symbol selection). Let i denote the step of  $\mathcal{T}_D$  and g(i) denote the step of M being simulated at step i of the decoder. At each decoding step we want to maintain the current Turing

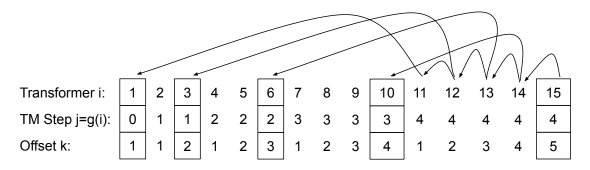


Figure 3: Mapping between transformer step and original Turing machine step.

machine state  $q^{g(i)}$  and symbol under the  $s^{g(i)}$  in  $\mathbf{y}_i$ . For roughly  $O(\sqrt{i})$  intermediate steps the state will remain the same, while we aggregate information about relevant past output symbols through sparse attention. To maintain the same state for intermediate steps, we introduce an extra switching layer (App. B.2.3). Finally, at the next compute step we will make the transition to new state  $q^{g(i)+1}$ , new head movement  $m^{g(i)}$ , and new output symbol  $v^{g(i)}$  to be written. Thereby we are able to completely simulate the given Turing machine M. As a result, we can prove the following main theorem:

**Theorem 3.** There exists a sparse attention mechanism using O(n) inner products such that the resulting class of Transformer Networks using this sparse attention mechanism is Turing Complete.

### Encoder

As [73], we use the same trivial single layer encoder where resulting  $K^{(e)}$  contains position embedding and  $V^{(e)}$  contains one-hot symbol representation.

### DECODER

Sparse Self-Attention mechanism for Decoder In this section, we will consider a particular instance of the sparse graph D at decoder. We define its edges to be given by the following relations:  $\forall j \in \mathbb{N}_+, 1 \leq k \leq j+1$ ,

$$\left(\frac{j(j+1)}{2}+k, \frac{k(k+1)}{2}\right)$$
 and

$$\left(\frac{j(j+1)}{2} + k, \frac{j(j+1)}{2} + k\right) \text{ if } k > 1 \text{ else } \left(\frac{j(j+1)}{2} + 1, \frac{j(j+1)}{2}\right).$$

This graph can be seen as a special case of BigBird where first type of edges are realizations of random and second type of edges correspond to locality. Also note that this graph satisfies the left-to-right constraint of decoder, i.e. no node attends to a node in the future.

**Embeddings and positional encodings** Our construction needs a different positional encoding  $pos_{Dec} : \mathbb{N} \to \mathbb{Q}^d$  for decoder:

$$\begin{array}{rcl} \mathrm{pos}_{\mathsf{Dec}}(i) & = & [ & 0, \dots, 0, \\ & & 0, \dots, 0, \\ & & 0, \dots, 0, \\ & & 1, g(i) + 1, \frac{1}{g(i) + 1}, \frac{1}{(g(i) + 1)^2}, h(i), 0, 0, 0, 0 & ] \end{array}$$

where  $g(i) = \left\lfloor \frac{-1+\sqrt{1+8i}}{2} \right\rfloor$  and h(i) = g(i+1) - g(i). Note that h(i) reduces to a binary indicator variable  $\mathbf{1}\left\{\frac{-1+\sqrt{1+8i}}{2} = \left\lfloor \frac{-1+\sqrt{1+8i}}{2} \right\rfloor \right\}$ .

# Induction Setup

We next show how to construct the decoder layers to produce the sequence of outputs  $y_1, y_2, ...$ , where  $y_i$  is given by:

$$\mathbf{y}_{i} = \begin{bmatrix} q^{g(i)} , q^{g(i)} , c^{g(i)} , c^{g(i)} , \\ 0, \dots, 0, \\ \mathbf{0}_{s}, 0, q^{g(i)} , \\ 0, 0, 0, 0, 0, u_{1}^{(i)}, u_{2}^{(i)}, u_{3}^{(i)}, u_{4}^{(i)} \end{bmatrix}$$

That is, at step i of our sparse decoder  $y_i$ , it will contain the information about the state of the turing machine M at time g(i), the symbol under the head of M at time g(i), and the current location of head of M at time g(i). We also have a placeholder symbol w and placeholder scalars  $u_1, u_2, u_3$ , whose role will be clear from our construction.

We consider as the starting vector for the decoder the vector

$$y_1 = [ [q_{\text{init}}], [\#], 0, \\ 0, \dots, 0, \\ 0, \dots, 0, \\ 0, \dots, 0$$

We assume that the start head is at  $c^{(0)} = 0$ , the initial state is  $q^{(0)} = q_{\text{init}}$ , and  $s^{(0)} = \#$  as we initialize from clean tape. We show the correctness of our construction by an inductive argument: we describe the architecture piece by piece and at the same time will show for every  $r \geq 0$ , our architecture constructs  $y_{r+1}$  from the previous vectors  $(y_0, \ldots, y_r)$ .

Thus, assume that  $y_1, \ldots, y_r$  satisfy the properties stated above. Since we are using positional encodings, the actual input for the first layer of the decoder is the sequence

$$y_1 + pos_{Dec}(1), y_2 + pos_{Dec}(2), ..., y_r + pos_{Dec}(r).$$

We denote by  $\overline{y}_i$  the vector  $y_i$  plus its positional encoding. Thus we have  $\forall 1 \leq i \leq r$  that

$$\begin{split} \overline{\boldsymbol{y}}_i &= & [ & \| \ q^{g(i)} \ \|, \| \ s^{g(i)} \ \|, c^{g(i)}, \\ & 0, \dots, 0, \\ & \boldsymbol{0}_s, 0, \| \ w^{(i)} \ \|, \\ & 1, g(i) + 1, \frac{1}{g(i) + 1}, \frac{1}{(g(i) + 1)^2}, h(i), u_1^{(i)}, u_2^{(i)}, u_3^{(i)}, u_4^{(i)} \ ] \end{split}$$

# B.2.1 Layer 1: Simulate Transition Function

In this layer, we use the cross-attention between encoder and decoder to access the input string and a feed-forward network to simulate the transition function of M. The first self attention in Eq. (5) is not used in this layer and we just produce the identity. This identity function is achieved by setting all queries, keys, values to be 0 everywhere plus the residual connection. Thus, we have  $p_i^1 = \overline{y}_i$ .

Since  $p_i^1$  is of the form  $[\_, \ldots, \_, 1, g(i) + 1, \_, \ldots, \_]$ , we know by Lemma B.1 of Pérez et al. [73] that if we use  $p_i^1$  to attend over the encoder we obtain

where  $\alpha$  and  $\beta$  are as defined in Eq. (21) of [73]. Thus in Eq. (4) we finally produce the vector  $\mathbf{a}_i^1$  given by

$$\mathbf{a}_{i}^{1} = \operatorname{CROSSATTN}(\mathbf{p}_{i}^{1}, \mathbf{K}^{e}, \mathbf{V}^{e}) + \mathbf{p}_{i}^{1} \\
= [ \mathbf{q}^{g(i)} ], [ \mathbf{s}^{g(i)} ], c^{g(i)}, \\
0, \dots, 0, \\
[ \alpha^{g(i)+1} ], \beta^{g(i)+1}, [ \mathbf{w}^{(i)} ], \\
1, g(i) + 1, \frac{1}{g(i)+1}, \frac{1}{(g(i)+1)^{2}}, h(i), u_{1}^{(i)}, u_{2}^{(i)}, u_{3}^{(i)}, u_{4}^{(i)} ]$$
(7)

As the final piece of the first decoder layer we use a function  $O_1(\cdot)$  (Eq. (3)) that satisfies the following lemma.

**Lemma 6** (Lemma B.2 [73]). There exists a two-layer feed-forward network  $O_1: \mathbb{Q}^d \to \mathbb{Q}^d$  such that with input vector  $\mathbf{a}_i^1$  (Eq. (7)) produces as output

$$\begin{array}{rcl} O_1(\boldsymbol{a}_i^1) & = & [ & 0, \dots, 0, \\ & & \mathbb{I} \ q^{g(i)+1} \ \mathbb{J}, \mathbb{I} \ v^{g(i)} \ \mathbb{J}, m^{g(i)}, 0, 0, 0, 0 \\ & & 0, \dots, 0, \\ & & 0, \dots, 0 \end{array} \right]$$

That is, function  $O_1(\cdot)$  simulates transition  $\delta(q^{g(i)}, s^{g(i)})$  to construct  $[q^{g(i)+1}]$ ,  $[v^{g(i)}]$ , and  $m^{g(i)}$  besides some other linear transformations.

Thus, finally the output of the first decoder layer is

## B.2.2 Layer 2: Finding Head Node

In this layer, we only use the feed-forward network to evaluate the next location of the head. The self-attention and cross-attention are set to be the identity function, so  $\boldsymbol{a}_i^2 = \boldsymbol{p}_i^2 = \boldsymbol{z}_i^1$ . Recall that  $c^{g(i)}$  is the cell to which M is pointing to at time g(i), and that it satisfies the following recursion  $c^{g(i)+1} = c^{g(i)} + m^{g(i)}$ , which can be expanded to see that that  $c^{g(i)+1} = m^{(0)} + m^{(1)} + \cdots + m^{g(i)}$ . Its not difficult to see that a two layer network with non-linearity can compute  $c^{g(i)+1}/(g(i)+1)$  and  $c^{g(i)}/(g(i)+1)$  from  $c^{g(i)}$ ,  $m^{g(i)}$ , and 1/(g(i)+1)

using the relation  $c^{g(i)+1} = c^{g(i)} + m^{g(i)}$ . At the end of layer 2, we obtain

### B.2.3 Layer 3: Distinguishing Node Type

This is an additional layer (not present in the work of [73]), where we propagate computations in our sparse graph. In particular, we will use this layer to "compute" or accumulate state in intermediate nodes. We make this clear below. The self-attention and cross-attention are all set to be the identity function, so  $a_i^3 = p_i^3 = z_i^2$ . In this layer, we only use the dense attention layers to select the newly computed states or to continue with previous states. Using idea similar to Lemma B.6 of [73], we can construct a dense network such that

$$O([x, y, z, b])) = \begin{cases} [\mathbf{0}, \mathbf{0}, \mathbf{0}, 0] & \text{if } b = 1, \\ [\mathbf{0}, z - y, -z, 0] & \text{if } b = 0. \end{cases}$$

The negatives are generated to offset results from skip connection. We utilize such network to switch Turing machine state and position embedding for intermediate steps to the values received from previous time step and do nothing for compute nodes. We use h(i) as the flipping bit b. Thus, at end of layer 3, we obtain

$$\mathbf{z}_{i}^{3} = O_{3}(\mathbf{a}_{i}^{3}) + \mathbf{a}_{i}^{3} = \begin{bmatrix} 0, \dots, 0, \\ \| \hat{q}^{(i)} \|, \| \hat{v}^{(i)} \|, \hat{c}^{(i)}, \frac{1}{g(i)+1}, \frac{1}{(g(i)+1)^{2}}, \frac{c^{g(i)+1}}{g(i)+1}, \hat{u}_{4}^{(i)}, \\ \| \hat{\alpha}^{(i)} \|, \hat{\beta}^{(i)}, \mathbf{0}_{s}, \\ 1, \hat{u}_{1}^{(i)}, \hat{u}_{2}^{(i)}, \hat{u}_{3}^{(i)}, h(i), 0, 0, 0, 0 \end{bmatrix}$$

where we used h(i) for selecting old states. In particular,

• We copy the input state and head position as is for intermediate nodes. We do not need to transition to next Turing machine states in these nodes.

$$\hat{q}^{(i)} = \begin{cases} q^{g(i)+1} & \text{if } h(i) = 1 \\ q^{g(i)} & \text{if } h(i) = 0 \end{cases}, \quad \hat{v}^{(i)} = \begin{cases} v^{g(i)} & \text{if } h(i) = 1 \\ w^{(i)} & \text{if } h(i) = 0 \end{cases}, \quad \hat{c}^{(i)} = \begin{cases} c^{g(i)+1} & \text{if } h(i) = 1 \\ c^{g(i)} & \text{if } h(i) = 0 \end{cases}.$$

• To preserve the symbol under the head for intermediate nodes, we copy the previous symbol to  $\alpha$  location and set  $\beta = g(i) + 1$ , as the symbol at  $\alpha$  location will be copied as the symbol under head for next transformer step by the final transformation layer if  $\beta = g(i) + 1$ . Thus, we correctly preserve the previous symbol under head as Turing machine does not transition these nodes. For compute nodes, things happen as usual.

$$\hat{\alpha}^{(i)} = \begin{cases} \alpha^{g(i)+1} & \text{if } h(i) = 1\\ s^{g(i)} & \text{if } h(i) = 0 \end{cases}, \qquad \hat{\beta}^{(i)} = \begin{cases} \beta^{g(i)+1} & \text{if } h(i) = 1\\ g(i)+1 & \text{if } h(i) = 0 \end{cases}.$$

• Finally for the intermediate nodes, we copy the position embedding corresponding to current best symbol w, which is stored in  $u_1, u_2, u_3, u_4$ . For compute node, we let the position embedding correspond to current Turing machine step.

$$\hat{u}_1^{(i)} = \begin{cases} g(i)+1 & \text{if } h(i)=1 \\ u_1^{(i)} & \text{if } h(i)=0 \end{cases}, \qquad \hat{u}_2^{(i)} = \begin{cases} \frac{1}{(g(i)+1)} & \text{if } h(i)=1 \\ u_2^{(i)} & \text{if } h(i)=0 \end{cases},$$
 
$$\hat{u}_3^{(i)} = \begin{cases} \frac{1}{(g(i)+1)^2} & \text{if } h(i)=1 \\ u_3^{(i)} & \text{if } h(i)=0 \end{cases}, \qquad \hat{u}_4^{(i)} = \begin{cases} \frac{c^{g(i)}}{g(i)+1} & \text{if } h(i)=1 \\ u_4^{(i)} & \text{if } h(i)=0 \end{cases}.$$

For further simplification note that g(i+1) = g(i) if h(i) = 0 else g(i) + 1 when h(i) = 1. With this fact, we can conclude that  $\hat{q}^{(i)} = q^{g(i+1)}$  and  $\hat{c}^{(i)} = c^{g(i+1)}$ . Thus, we can write,

### B.2.4 Layer 4: Finding next symbol on tape

To find the symbol on tape under next head position  $c^{g(i)+1}$ , we try to find what was written last at the location  $c^{g(i)+1}$ . To facilitate this, following [73], we define  $\ell(j)$  to be the last time (previous to j) in which M was pointing to position  $c^{(j)}$ , or it is j-1 if this is the first time that M is pointing to  $c^{(j)}$ . Recall j is the Turing machine step counter, which is different from sparse transformer step i. [73] could utilize full attention mechanism to find  $v^{\ell(j+1)}$  at one go, but we have to do it over multiple steps owing to our sparse attention mechanism.

We use similar query, key, value functions as used for full attention by [73]  $\forall i$ :

$$\begin{array}{lll} Q_4(\boldsymbol{z}_i^3) & = & [ & 0, \dots, 0 \\ & & 0, \dots, 0, \\ & & 0, \dots, 0, \\ & & 0, \frac{c^{g(i)+1}}{g(i)+1}, \frac{1}{g(i)+1}, \frac{1}{3(g(i)+1)^2}, 0, 0, 0, 0, 0 ] \\ K_4(\boldsymbol{z}_i^3) & = & [ & 0, \dots, 0 \\ & & 0, \dots, 0, \\ & & 0, \dots, 0, \\ & & & 0, \hat{u}_2^{(i)}, \hat{u}_4^{(i)}, \hat{u}_3^{(i)}, 0, 0, 0, 0, 0, 0 ] \\ V_4(\boldsymbol{z}_i^3) & = & [ & 0, \dots, 0, \\ & & & 0, 0, 0, 0, 0, \hat{u}_1^{(i)}, \hat{u}_2^{(i)}, \hat{u}_3^{(i)}, \hat{u}_4^{(i)} & ] \\ \end{array}$$

It is clear that the three functions are linear transformations and thus they can be defined by feed-forward networks. Notice that the query vector is always formed using current time step position embedding, whereas key and value vectors are formed using copied over entries for intermediate nodes and using current entries only for compute node. Pérez et al. [73] find the desired  $v^{l(j+1)}$  as  $v^{m(j)}$  using full attention, where

$$m(t) = \underset{m \in \{0,...,t\}}{\operatorname{arg \, min}} \chi_t^j = \underset{m \in \{0,...,t\}}{\operatorname{arg \, min}} |\langle Q_4(\boldsymbol{z}_j^3), K_4(\boldsymbol{z}_m^3) \rangle|$$

Note the minimization is only over Turing machine steps, i.e. over compute nodes in our case. We show below that we can estimates m(j) by parts using sparse attention mechanism. The main idea is just to notice that minimization problem  $\min_{m \in \{0,...,t\}} \chi_t^j$  can be expressed as  $\min\{\cdots \min\{\min\{\chi_0^j,\chi_1^j\},\chi_2^j\},...,\chi_t^j\}$  by the associativity property.

By definition of our graph D, at every intermediate node i of the form j(j+1)/2 + k, i.e. where k > 0, g(i) = j and h(i) = 0, we will attend over node k(k+1)/2 and best till now copied from i-1. The node k(k+1)/2 is never an intermediate node as h(k(k+1)/2) = 1 for all k and in fact corresponds to Turing machine's step k. This will help us select the key and value corresponding to min between node k(k+1)/2 and i-1. In other words, at node i of the form j(j+1)/2 + k we would have evaluated m(k) and corresponding value selected:

$$w^{(j(j+1)/2+k+1)} = \hat{v}^{m(k-1)}$$

and similarly for u's. So after going through all the intermediate nodes, finally at the next compute node, i.e. when k = j + 1, we will obtain the minimum value over all of 0, 1, ..., j. This implies at a compute node will be able to recover  $\ell(g(i) + 1)$  and its corresponding value as shown in Lemma B.4 of [73]. Then we have that  $p_i^4$  is given by

$$\begin{aligned} \boldsymbol{p}_{i}^{4} &= & \operatorname{ATTN}_{D}(\boldsymbol{Z}_{i}^{3}) + \boldsymbol{z}_{i}^{3} \\ &= & [ & 0, \dots, 0, \\ & & & \| q^{g(i+1)} \|, \| \hat{\boldsymbol{v}}^{(i)} \|, c^{g(i+1)}, 0, \frac{c^{g(i)+1}}{g(i)+1}, \hat{\boldsymbol{u}}_{4}^{(i)}, \\ & & & \| \hat{\boldsymbol{\alpha}}^{(i)} \|, \hat{\boldsymbol{\beta}}^{(i)}, \| \boldsymbol{w}^{(i+1)} \|, \\ & & 1, \hat{\boldsymbol{u}}_{1}^{(i)}, \hat{\boldsymbol{u}}_{2}^{(i)}, \hat{\boldsymbol{u}}_{3}^{(i)}, h(i), \boldsymbol{u}_{1}^{(i+1)}, \boldsymbol{u}_{2}^{(i+1)}, \boldsymbol{u}_{3}^{(i+1)}, \boldsymbol{u}_{4}^{(i+1)} \end{bmatrix} \end{aligned}$$
(8)

The cross-attention and feed-forward network are set to be identity, so  $z_i^4 = a_i^4 = p_i^4$ .

#### B.2.5 Final transformation

We finish our construction by using the final transformation function  $F(\cdot)$  from the corresponding lemma from Pérez et al. [73], with a slight modification.

**Lemma 7** (Lemma B.5 [73]). There exists a function  $F: \mathbb{Q}^d \to \mathbb{Q}^d$  defined by a feed-forward network such that

$$\begin{split} F(\boldsymbol{z}_{r}^{4}) &= & [ & \llbracket \ q^{g(r+1)} \ \rrbracket, \llbracket \ s^{g(r+1))} \ \rrbracket, c^{g(r+1)}, \\ & 0, \dots, 0, \\ & \boldsymbol{0}_{s}, 0, \llbracket \ w^{(r+1)} \ \rrbracket, \\ & 0, 0, 0, 0, 0, u_{1}^{(r+1)}, u_{2}^{(r+1)}, u_{3}^{(r+1)}, u_{4}^{(r+1)} ] \\ &= & \boldsymbol{u}_{r+1} \end{split}$$

The modification is to let  $w, u_1, u_2, u_3$  to pass through. This yields the desired input to transformer at next time step for both intermediate and compute node, thereby concluding our induction.

# Appendix C. Limitations

Finally, we show that sparse attention mechanisms can not universally replace dense attention mechanisms, i.e. there is no free lunch. We demonstrate a natural task which can be solved by the full attention mechanism in O(1)-layers. However, under standard complexity theoretic assumptions, we show that this problem will require  $\tilde{\Omega}(n)$ -layers for any sparse attention layers with  $\tilde{O}(n)$  edges (not just BigBird). (We use the standard notation  $\tilde{\Omega}(n)$  to hide the dependence on poly-logarithmic factors.)

We consider the simple problem of finding the corresponding furthest vector for each vector in the given sequence of length n and dimension  $d \in \Theta(\log^2 n)$ . The assumption on the dimension is mild, as in many situations the dimension d = 768 is actually comparable to the number of n.

**Task 1.** Given 
$$n$$
 unit vectors  $\{u_1, \ldots, u_n\}$ , each in  $\mathbb{R}^d$  where  $d = \Theta(\log^2 n)$ , find  $f(u_1, \ldots, u_n) \to (u_{1^*}, \ldots, u_{n^*})$  where for a fixed  $j \in [n]$ , we define  $j^* = \arg \max_k \|u_k - u_j\|_2^2$ .

Finding vectors that are furthest apart boils down to minimizing inner product search in case of unit vectors. For a full-attention mechanism with appropriate query and keys, this task is very easy as we can evaluate all pair-wise inner products.

The impossibility for sparse-attention follows from hardness results stemming from Orthogonal Vector Conjecture (OVC) [2, 1, 97, 7]. The OVC is a widely used assumption in fine-grained complexity. Informally, it states that one cannot determine if the minimum inner product among n Boolean vectors is 0 in subquadratic time.

Conjecture 1 (Orthogonal Vectors Conjecture). For every  $\epsilon > 0$ , there is a  $c \ge 1$  such that given n Boolean vectors in d dimension, cannot determine if there is a pair of orthogonal vectors in  $O(n^{2-\epsilon})$  time on instances with  $d \ge c \log n$ .

We show a reduction using conjecture 1 to show that a transformer  $g \in \mathcal{T}_D^{H=O(d),m=O(d),q=O(d)}$  for any sparse directed graph D which completes Task 1 must require a superlinear number of layers.

**Proposition 2.** There exists a single layer full-attention network  $g \in \mathcal{T}^{H=1,m=2d,q=0}$  that can evaluate Task 1, i.e.  $g(u_1,...,u_n) = [u_{1^*},...,u_{n^*}]$ , but for any sparse-attention network in  $\mathcal{T}_D^{H=O(d),m=O(d),q=O(d)}$  with graph D having  $\tilde{O}(n)$  edges (i.e. inner product evaluations), would require  $\tilde{\Omega}(n^{1-o(1)})$  layers.

*Proof.* We will break this proof into two parts:

Part 1: The full attention mechanism can solve the problem in O(1) layer We begin by providing an explicit construction of a single layer full self-attention that can evaluate Task 1.

**Step 1** We embed each  $u_i$  in the input into  $\mathbb{R}^{2d}$  as follows:

$$x_i := E(u_i) = [u_i; 0] \tag{9}$$

**Step 2** Construct query, key, value functions as follows:

$$Q([a;b]) = -a$$
  
 $K([a;b]) = a$  (10)  
 $V([a;b]) = [0;a]$ 

Then Attn $(Q(x_i), K(X), V(X)) = [0; u_{\arg\max_i \langle -u_i, u_i \rangle}]$ . Then,

$$a_i = \text{Attn}(Q(x_i), K(X), V(X)) + x_i = [u_i; u_{\arg\max_i \langle -u_i, u_i \rangle}] = [u_i; u_{i^*}]$$
 (11)

**Step 3** Let  $O(a_i) = 0$ , then the output  $z_i = [u_i; u_{i*}]$  as desired.

To complete the argument, observe that it now only takes O(n) inner products to check if there is a pair of orthogonal vectors as we need only compare  $\langle u_i, u_{i^*} \rangle$ .

Part 2: Every Sparse Attention Mechanism will need  $\tilde{\Omega}(n^{1-\epsilon})$  layers We prove by contradiction that it is impossible to solve Task 1 by any  $g \in \mathcal{T}_D^{H=O(d),m=O(d),q=O(d)}$  sparse-attention graph D with  $\tilde{O}(n)$  edges.

Suppose we can solve Task 1 using a network  $g \in \mathcal{T}_D^{H=O(d), m=O(d), q=O(d)}$  that has l layers. Recall that all the computation we do in one layer is:

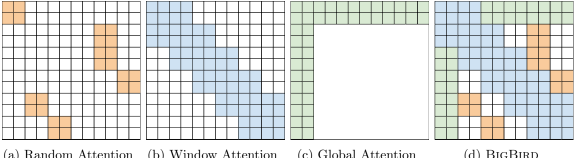
$$a_i = \text{Attn}_D(Q(x_i), K(X_{N(i)}), V(X_{N(i)}) + x_i$$
  
 $x_i = O(a_i) + a_i$  (12)

where  $Attn_D$  is defined in eq. (AT).

Thus, total computation per layer is  $\tilde{O}(nd^3)$  and consequently  $\tilde{O}(nld^3)$  for the whole network consisting of l layers.

We can use the result of Task 1 to solve the orthogonal vector (OV) problem (defined in Conjecture 1) in linear time. So in total, we will be able to solve any instance of OV in  $\tilde{O}(nld^3)$  time.

Now if  $l = O(n^{1-\epsilon})$  for any  $\epsilon > 0$  and  $d = \Theta(\log^2 n)$ , then it appears that we are able to solve OV in  $\tilde{O}(n^{2-\epsilon})$  which contradicts Conjecture 1. Therefore, we need at least  $\tilde{\Omega}(n^{1-o(1)})$  layers.



(a) Random Attention (b) Window Attention

(c) Global Attention

(d) BigBird

Figure 4: Building blocks of the block-attention mechanism used in BIGBIRD with block size = 2. This implies the attention matrix is split into blocks of size  $2 \times 2$ . All the previous BIGBIRD parameters work on each block as a unit. White color indicates absence of attention. (a) random attention with r=1, (b) sliding window attention with w=3 (c) global attention with q = 1. (d) the combined BigBird model.

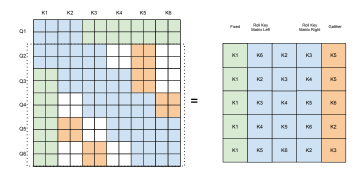


Figure 5: Structured block sparsity in BIGBIRD helps in compactly packing our operations of sparse attention, thereby making our method efficient on GPU/TPU. The matrix on the left is same as that shown in Fig. 4d. The figure on the right shows how the attention is actually calculated for all blocks of query except the first. The green column is fixed, the second column can be obtained by rotating the key matrix to the left, the third column is the key matrix, the fourth column can be obtained by rotating the key matrix right, and then only for the last column we need to use the computationally inefficient gather operation. Even then we do block gather rather than gathering keys separately for each query, further speeding up operations in the GPU/TPU.

## Appendix D. Implementation details

We have optimized the code for modern hardware. Hardware accelerators like GPUs and TPUs truly shine on coalesced memory operations which load blocks of contiguous bytes at once. Thus, its not very efficient to have small sporadic look-ups caused by sliding window or a random elements. We alleviating this by "blockifying" the entries.

**GPU/TPU** and **Sparsity** Ideally, if the adjacency matrix A described in Sec. 2 is sparse, one would hope this would be sufficient to speed up the implementation. Unfortunately, it is

well known [34, 103], that such sparse multiplications cannot be efficiently implemented in GPUs. GPUs have thousands of cores performing operations in parallel. Thus, we would not be able to efficiently perform the sparse matrix multiplication mentioned in section Sec. 2.

As a result we propose to first blockify the Attention matrix i.e. we pack sets of query and keys together and then define attention on these blocks. It is easier to explain this process using the example shown in Fig. 4. Suppose, there are 12 query and 12 key vectors to attend to. Using a block size of 2, we split the query matrix into 12/2 = 6 blocks and similarly the key matrix into 12/2 = 6 blocks. Then the three different building components of BIGBIRD are defined on the block matrix. In particular the three different components are:

- 1. Random Attention: Each query block attends to r random Key blocks. In Fig. 4a, r=1 with block size 2. This implies that each query block of size 2 randomly attends to a key block of size 2.
- 2. Window Attention: While creating the block, we ensure that the number of query blocks and the number of key blocks are the same. This helps us in defining the block window attention. Every query block with index j attends to key block with index j (w-1)/2 to j + (w-1)/2, including key block j. In Fig. 4b, w = 3 with block size 2. It means that each query block j (size 2 queries) attends to key block j 1, j, j + 1.
- 3. Global attention: Global attention remains the same as defined in Sec. 2, but we compute it in terms of blocks. In Fig. 4c, g = 1 with block size 2. For BIGBIRD-ITC this implies that one query and key block, attend to everyone.

The resulting global attention matrix is shown in Fig. 4d.

We use another trick to speed-up computation. For multiplying pairs of vectors with indexes in a matrix in a GPU, we need to use gather operation. Unfortunately this operation is not efficient and one would want to reduce its use as much as possible. Upon closer examination of window and global attention, we observe that we can perform them without using a gather operation. The global attention for all blocks other than Q1 in Fig. 5, can be computed if we "roll" the key-block matrix forward and backward and multiply with the query block matrix as shown in Fig. 5. While rolling, we do not include global key blocks (K1 in Fig. 5). We only use gather for the random attention, which is very small (r=3 for all of our experiments). This improves our efficiency. As seen in Fig. 5, rolling of window attention, leads to Q2 query block attending to K6 key block and Q6 attending to K2, which were not present in the original attention.

# Appendix E. NLP experiments details

### E.1 MLM Pretraining

We use four publicly available datasets Books [110], CC-News [35], Stories [90] and Wikipedia to pretrain BIGBIRD. We borrow the sentencepiece vocabulary as RoBERTa (which is in turn borrowed from GPT2). We split any document longer than 4096 into multiple documents and we joined multiple documents that were much smaller than 4096. Following the original BERT training, we mask 15% of tokens in these four datasets, and train to predict the mask. We warm start from RoBERTa's checkpoint. We train two different models: BIGBIRD-ITC-base and BIGBIRD-ETC-base. The hyper-parameters for these two models are given in Tab. 12. In all experiments we use a learning rate warmup over the first 10,000 steps, and linear decay of the learning rate.

Similar to the norm, we trained a large version of model as well, which has 24 layers with 16 heads and hidden dimension of 1024. Following the observation from RoBERTa, we pretrain on a larger batch size of 2048 for this size. For BIGBIRD-ITC the block length was kept same as base size, but for BIGBIRD-ETC the block length was almost doubled to 169. All the remaining parameters were the same.

Parameter	BigBird-itc	BIGBIRD-ETC
Block length, b	64	84
# of global token, $g$	$2 \times b$	256
Window length, $w$	$3 \times b$	$3 \times b$
# of random token, $r$	$3 \times b$	0
Max. sequence length	4096	4096
# of heads	12	12
# of hidden layers	12	12
Hidden layer size	768	768
Batch size	256	256
Loss	MLM	MLM
Activation layer	gelu	gelu
Dropout prob	0.1	0.1
Attention dropout prob	0.1	0.1
Optimizer	Adam	Adam
Learning rate	$10^{-4}$	$10^{-4}$
Compute resources	$8 \times 8 \text{ TPUv3}$	$8 \times 8 \text{ TPUv3}$

Table 12: Hyperparameters for the two BigBird base models for MLM.

# E.2 Question Answering

The detailed statistics of the four datasets used are given in Tab. 13. All the hyperparameters for BigBird, used for creating Tab. 4 are shown in Tab. 14 and those submitted to get Tab. 5 are shown in Tab. 15. We use two types of regularization in training:

- We used a variant of contrastive predictive coding [71] as a dual encoder model.
- We use position embedding for ITC and relative position encoding [80] for ETC.

Next, we will mention the dataset/task specific part of the model.

	Instan	ces	Instance 1	Length
Dataset	Training	Dev	Median	Max
HotpotQA-distractor [101]	90447	7405	1227	3560
Natural Questions [53]	307373	7830	3258	77962
TriviaQA [42]	61888	7993	4900	32755
WikiHop [96]	43738	5129	1541	20337

Table 13: Question Answering Datasets

Parameter	Hotp	$\operatorname{HotpotQA}$		NaturalQ		TriviaQA		WikiHop	
Global token location	ITC	ETC	ITC	ETC	ITC	ETC	ITC	ETC	
# of global token, $g$	128	256	128	230	128	320	128	430	
Window length, $w$	192	252	192	252	192	252	192	252	
# of random token, $r$	192	0	192	0	192	0	192	0	
Max. sequence length	4096	4096	4096	4096	4096	4096	4096	4096	
# of heads	12	12	12	12	12	12	12	12	
# of hidden layers	12	12	12	12	12	12	12	12	
Hidden layer size	768	768	768	768	768	768	768	768	
Batch size	32	32	128	128	32	32	64	64	
T	cross-e	ntropy	cross-e	ntropy	cross-€	entropy	cross-er	ntropy	
Loss	golden	spans	golden	golden spans		noisy spans [19]		ans choices	
Compute resources	$4 \times 2$ [	$\Gamma PUv3$	$4 \times 8$ 7	$\Gamma PUv3$	$4 \times 2$	TPUv3	$4 \times 4$ T	PUv3	

Table 14: Hyperparameters of base BIGBIRD model used for Question Answering i.e. the numbers reported in Tab. 4

HotpotQA The data consists of each question with multiple evidence paragraphs. We filtered 16 QA where the answer was not in the given evidences. For BIGBIRD-ITC, we use first 128 global tokens. For BIGBIRD-ETC, we have one global token for each question token, one for each evidence paragraph, and one for each sentence within the paragraph, for a maximum of 256 global token. We use a dense layer on the output corresponding to global token of the evidence paragraph to predict whether its a supporting fact with a threshold over the output logits. The answer type (yes/no/span) is predicted with a single dense layer from the global CLS token. For span based answers, the spans are predicted with dense layers on the sequence with the distance between start and end positions to be no more than 30 words. The spans are ranked by sum of start and end logits.

Natural Questions Here also the data consists of question with supporting evidence, but in form of a single, potentially long, document and not multiple paragraphs. We largely follow the setup of [5]. For documents, that are longer than 4096, a sliding window approach is used with stride of 2048. We use CLS token at the beginning, followed by the question followed by a separator token followed by the document as input. For BIGBIRD-ITC, we make the first 128 tokens as global. For BIGBIRD-ETC, we make a global token for CLS, question, and one token for each of the paragraphs. We train four predictors at the final layer to predict long answer start, long answer end, short answer start and short answer end

Parameter	HotpotQA	NaturalQ	TriviaQA	WikiHop
Global token location	ETC	ETC	ETC	ETC
# of global token, $g$	256	230	320	430
Window length, $w$	507	507	507	507
# of random token, $r$	0	0	0	0
Max. sequence length	4096	4096	4096	4096
# of heads	16	16	16	16
# of hidden layers	24	24	24	24
Hidden layer size	1024	1024	1024	1024
Batch size	32	64	32	64
Loss	cross-entropy	cross-entropy	cross-entropy	cross-entropy
Num epochs	$\{5, 9\}$	$\{3, 5\}$	$\{4, 6\}$	$\{5, 10\}$
Optimizer	Adam	Adam	Adam	LAMB
Learning rate	$3 \times 10^{-5}$	$\{5,10\} \times 10^{-5}$	$\{3,5\} \times 10^{-5}$	$\{2,5\} \times 10^{-5}$
Compute resources	$4\times4~\mathrm{TPUv3}$	$4 \times 8 \text{ TPUv3}$	$4 \times 4$ TPUv3	$4 \times 8 \text{ TPUv3}$

Table 15: Hyperparameters of large BigBird model for Question Answering submitted for test i.e. the numbers reported in Tab. 5

respectively. Instead of independently predicting the start and end of answers we first predict the start and then predict the best end location beyond the start. For short answer, we limit the distance between start and end positions to be no more than 38 words. The answer type (null, yes, no, short, long) is predicted from CLS token output embedding. When the logit for a yes/no answer is higher than the logits for short, long or null answer, we replace the short answer with a corresponding yes/no text.

TriviaQA The data consists of question-answer pairs with Wikipedia articles as the "noisy" supporting evidence. We call them noisy because the given Wikipedia articles may or may not contain the answer. Moreover, the answer entities is not annotated to appropriate span in the article, rather all occurrences found using fuzzy string matching are listed. We use CLS token at the beginning, followed by the question followed by a separator token followed by the document as input. For BIGBIRD-ITC, we make the first 128 tokens as global. For BIGBIRD-ETC, we make a global token for CLS, question, and one token for each sentence up to a maximum of 320 global tokens. Given the noisy nature of answer span, we follow Clark and Gardner [19] for training. We use a dense layer on the sequence to predict the answer span for each article independently, with the distance between start and end positions to be no more than 16 words. For each article the span with maximum start logit + end logit is chosen. Then we normalize over all the documents associated with that question.

WikiHop For each question in WikiHop, we are given upto 79 candidates, and 63 supporting paragraphs. In our BigBird-itc model, following Beltagy et al. [8], we concatenate the answer and the question with special tokens, [q] Question [/q] [ans] Ans1 [/ans] ... [ans] AnsN [/ans] along with the context. As the start of the text, always contains questions followed by answers, we make the first 128 token attend globally. In BigBird-etc model, we do not need to insert special [ans], [/ans] etc. as we design global tokens appropriately. Along with global tokens for question, we have one per candidate answer up

to a maximum of 430. Further, we linked answer tokens to their mentions using relative position label. Lastly, we use a dense layer that takes in the output vector corresponding to a candidate answer, and predicts a score for the current candidate to be the correct answer. We apply this dense layer to each candidate independently and the candidate with the best score is picked as our final answer.

It is worthwhile to note that explicitly designed attention connection in ETC works slightly better, the random connection based ITC is pretty competative.

### E.3 Document Classification

**Experimental Setup** Tab. 16 gives the detail list of hyperparameters used to get results shown in Tab. 6. In all experiments we use a learning rate warmup over the first 10% steps, and linear decay of the learning rate.

Parameter	IMDb	Arxiv	Patents	Hyperpartisan	Yelp-5
Batch size	64	64	64	32	32
Learning rate	$1 \times 10^{-5}$	$3 \times 10^{-5}$	$5 \times 10^{-5}$	$5 \times 10^{-6}$	$2 \times 10^{-5}$
Num epochs	40	10	3	15	2
TPUv3 slice	$4 \times 4$	$4 \times 4$	$4 \times 4$	$4 \times 2$	$4 \times 8$
# of heads			12		16
# of hidden layers			12		24
Hidden layer size			768		1024
Block length, $b$			64		
Global token location			ITC		
# of global token, $g$			$2 \times b$		
Window length, $w$			$3 \times b$		
# of random token, $r$			$3 \times b$		
Max. sequence length			4096		
Vocab size			50358		
Activation layer			$\operatorname{gelu}$		
Dropout prob			0.1		
Attention dropout prob			0.1		
Loss			cross-entro	ору	
Optimizer			Adam		

Table 16: Hyperparameters for document classification.

System	MNLI-(m/mm) 392k	QQP 363k	QNLI 108k	SST-2 67k	CoLA 8.5k	STS-B 5.7k	MRPC 3.5k	RTE 2.5k
BERT	84.6/83.4	71.2	90.5	93.5	52.1	85.8	88.9	66.4
XLNet	86.8/-	91.4	91.7	94.7	60.2	89.5	88.2	74.0
RoBERTa	87.6/-	91.9	92.8	94.8	63.6	91.2	90.2	78.7
BigBird	87.5/87.3	88.6	92.2	94.6	58.5	87.8	91.5	75.0

Table 17: GLUE Dev results on base sized models. Number of training examples is reported below each task. MCC score is reported for CoLA, F1 score is reported for MRPC, Spearman correlation is reported for STS-B, and accuracy scores are reported for the other tasks.

**GLUE** The General Language Understanding Evaluation (GLUE) benchmark [93], test language models on 8 different natural language understanding tasks. We used the same training parameters as mentioned in https://github.com/pytorch/fairseq/blob/master/examples/roberta/README.glue.md. Our model parameters are  $b = 64, g = 2 \times b, w = 3 \times b, r = 3 \times b$  ( we used the BIGBIRD-ITC base model pretrained on MLM task). We compare the performance of BIGBIRD to BERT, XLNet [102] and RoBERTa in Tab. 17. We find that even on task that have a much smaller context, our performance is competitive to full attention models.

#### E.4 Summarization

As discussed in Sec. 4.2, given the small length of output sequence, we used sparse BIGBIRD attention only for encoder, while keeping the full attention for decoder. The number of hidden layers, number of heads, and hidden dimension is same for encoder and decoder. The hyperparameters are detailed in Tab. 18. We summarize our result in Tab. 20. In all experiments, we use a learning rate warmup over the first 10,000 steps, and square root decay of the learning rate.

Parameter	Base: BIGBIRD-R	oBERTa	Large: BIGBIRD-Pegasus
Block length, b		64	64
Global token location		ITC	ITC
# of global token, $g$		$2 \times b$	$2 \times b$
Window length, $w$		$3 \times b$	$3 \times b$
# of random token, $r$		$3 \times b$	$3 \times b$
	BBC-XSUM:	1024	1024
Max. encoder sequence length	CNN/DM:	2048	2048
	Others:	3072	3072
	BBC-XSUM:	64	64
Max. decoder sequence length	CNN/DM:	128	128
	Others:	256	256
Beam size		5	5
Length penalty	BBC-XSUM:	0.7	0.7
Length penalty	Others:	0.8	0.8
# of heads		12	16
# of hidden layers		12	16
Hidden layer size		768	1024
Batch size		128	128
Loss	teache	er forced	teacher forced
1055	cross	-entropy	cross-entropy
Activation layer		$\operatorname{gelu}$	gelu
Dropout prob		0.1	0.1
Attention dropout prob		0.1	0.1
Optimizer		Adam	Adafactor
Learning rate		$1 \times 10^{-5}$	$1 \times 10^{-4}$
Compute resources	$4 \times 4$	TPUv3	$4 \times 8 \text{ TPUv3}$

Table 18: Encoder hyperparameters for Summarization. We use full attention in decoder

	Ir	Instances		Input	Length	Output	Output Length	
Dataset	Training	Dev	Test	Median	90%-ile	Median	90%-ile	
BBC XSum [68]	204044	11332	11334	359	920	25	32	
CNN/DailyMail [37]	287113	13368	11490	777	1439	59	93	

Table 19: Shorter summarization dataset statistics.

		В	BC XSu	m	CN	$\mathrm{CNN}/\mathrm{DailyMail}$		
Model		R-1	R-2	R-L	R1	R2	R-L	
	Lead	16.30	1.61	11.95	39.60	17.70	36.20	
	PtGen [78]	29.70	9.21	23.24	39.53	17.28	36.38	
	ConvS2S [29]	31.89	11.54	25.75	_	_	_	
Art	MMN [49]	32.00	12.10	26.00	_	_	_	
	Bottom-Up [30]	_	_	_	41.22	18.68	38.34	
Prior	TransLM [46]	_	_	_	39.65	17.74	36.85	
щ	UniLM [24]	_	_	_	43.47	20.30	40.63	
	Extr-Abst-BERT [63]	38.81	16.50	31.27	42.13	19.60	39.18	
	BART [57]	45.14	22.27	37.25	44.16	21.28	40.90	
	Transformer [92]	29.61	9.47	23.17	34.89	13.13	32.12	
Ş.	+ RoBERTa [77]	39.92	<u>17.33</u>	32.63	39.44	18.69	36.80	
$\mathbf{Base}$	+ Pegasus [107]	39.79	16.58	31.70	41.79	<u>18.81</u>	38.93	
	BIGBIRD-RoBERTa	39.52	17.22	32.30	39.25	18.46	36.61	
е	Pegasus (Reported) [107]	47.60	24.83	39.64	44.16	21.56	41.30	
Large	Pegasus (Re-eval)	47.37	24.31	39.23	44.15	21.56	41.05	
La	BigBird-Pegasus	47.12	24.05	38.80	43.84	21.11	40.74	

Table 20: Summarization ROUGE score for shorter documents.

Following success of several recent works [77, 64], we warm start our encoder-decoder BIGBIRD transformer model with pretrained weights and the weights between encoder and decoder are shared. In particular, the query/key/value matrix of self-attention and all the feedforward layers are shared between encoder and decoder. The only variable that is initialized randomly is the encoder-decoder attention. For base sized model, we utilize our MLM pretrained model on 4096 sequence length from App. E.1, which is in turn initialized using the public RoBERTa checkpoint. For the large size model, we lift weight from the state-of-the-art Pegasus model [107], which is pretrained using an objective designed for summarization task.

To check if sparse attention causes significant degradation as compared to full attention, we further experiment on two shorter but popular datasets, where full attention can be used without significantly truncating the document. The statistics of these two datasets are in Tab. 19. We see that our performance is competitive, which shows that sparse attention can achieve similar performance to a full attention models.

# Appendix F. Genomics experiments details

In this section we provide details of the experimental setup for BIGBIRD on genomics data.

# F.1 Pretraining

We try to keep the experimental setup as close to a typical NLP pipeline. In this regard, we take human reference GRCh37<sup>6</sup> and convert it into documents  $\mathcal{D}$ . Each document  $d \in \mathcal{D}$  is a sequence of sentences, where each sentence is a sequence of fragments of DNA. We construct the documents as follows:

- 1. Start with empty document set  $D = \emptyset$ .
- 2. For each chromosome C, repeat the following procedure 10 times.
  - (a) Pick uniformly at random a starting point q between base pairs 0 and 5000 from the 5' end.
  - (b) Repeat until q > |C|
    - i. Pick uniformly at random s a number between 50 and 100 to denote number of sentences per document.
    - ii. Constructs a document d containing s sentences using consecutive base pairs (bps). The length of each sentence is chosen uniformly at random between 500-1000. Thus the resulting document has 25,000 100,000 bps.

iii. 
$$D = D \bigcup d$$

iv. 
$$q = q + |d|$$

By this procedure we end-up with approximately 450K documents.

Next we run sentencepiece [51] tokenization on the resulting documents. In particular, using 5 characters as the building blocks (four for bases - A, T, C, G and one for missing symbol N), we construct a byte pair encoding table of size 32k, with each token representing 8.78 base pairs on average.

Using the above constructed documents, we construct a dataset for two pretraining tasks following Devlin et al. [23]:

- Masked Language Model (MLM): In order to train a deep bidirectional representation, BERT training introduces the MLM task, where we simply mask out 15% of the input tokens at random, and then predict those masked tokens. We can simply replace such masked out of the tokens with a [MASK] placeholder, but it leads to a distribution mis-match for downstream tasks which will not have such placeholders. To mitigate with this issue, out of the 15% of the tokens selected for masking:
  - 80\% of the tokens are actually replaced with the token [MASK].

<sup>6.</sup> https://www.ncbi.nlm.nih.gov/assembly/GCF\_000001405.39

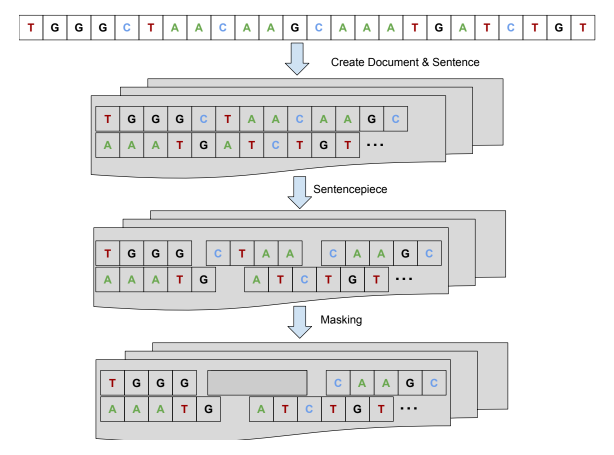


Figure 6: Visual description of how the masked language modeling data was generated from raw DNA dataset. The raw DNA sequences of GRCh37, where split at random positions to create documents with 50-100 sentences where each sentence was 500-1000 base pairs (bps). Thus each document had a continuous strand of 25000-100,000 bps of DNA. This process was repeated 10 times to create 10 sets of document for each chromosome of GRCH37. The resulting set of documents was then passed through Sentencepiece that created tokens of average 8bp. For pretraining we used masked language model and masked 15% of the tokens and trained on predicting the masked tokens.

- -10% of the time tokens are replaced with a random token.
- 10\% of the time tokens are left unchanged, but are still predicted at output.

We run this entire sequence through the BIGBIRD transformer encoder and then predict corresponding to the masked positions, based on the context provided by the other non-masked tokens in the sequence.

• Next Sentence Prediction (NSP): In order to understand relationship between two sequences, BERT training introduces the NSP task, where we predict if a given pair of sequences are contiguous or not. During training the model gets as input pairs of sequences separated by [SEP] token along with a [CLS] token at the start. Overall the input pattern is: [CLS] sequence A [SEP] sequence B [SEP]. For 50% of the time the second sequence comes from true sequence after the first one. Remaining 50% of the time it is a a random sequence from the full dataset. The model is then required to predict this relationship using the output corresponding to the [CLS] token, which is fed into a simple binary classification layer.

The sequence of steps is visually elaborated in Fig. 8. The model is trained with both MLM and NSP together. Training hyperparameter is provided in second columns of Tab. 21. In all experiments we use a learning rate warmup over the first 10,000 steps, and linear decay of the learning rate.

We additionally performed a simple ablation study to validate the hypothesis, that similar to NLP, having a larger context improves performance. We use MLM task described above to test how BigBird performed with sequences of different length. Accuracy on MLM task with increasing sequence length is shown in Fig. 7. Not only longer context improves final accuracy, it also leads to faster learning, as we have now more opportunities for masking.

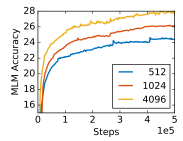


Figure 7: BIGBIRD accuracy with context length.

#### F.2 Promoter Region Prediction

The promoter region plays an important role in transcription initiation and thus its recognition is an important area of interest in the field of bioinformatics. Following Oubounyt et al. [72], we use datasets from Eukaryotic Promoter Database (EPDnew) [25], which contains 29,597 promoter region in the human genome. Around the transcription start site (TSS), we extract a sequence of 8000 bp (-5000 +3000 bp) from the human reference genome GRCh37. Since EPDnew uses newer GRCh38, we convert to GRCh37 coordinates using LiftOver [45].

Following Oubounyt et al. [72] for each promoter region example, a negative example (non-promoter sequences) with the same size of the positive one is constructed as follow: The positive sequence is divided into 20 subsequences. Then, 12 subsequences are picked randomly and substituted randomly. The remaining 8 subsequences are conserved. This process is illustrated in Figure 1 of [72]. Applying this process to the positive set results in new non-promoter sequences with conserved parts from promoter sequences (the unchanged

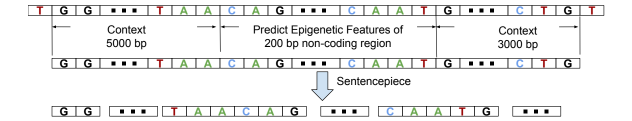


Figure 8: Visual description of the DNA segment from which we predict the chromatin profile for a given non-coding region of the raw DNA sequences of GRCh37. We take 8000 bps of DNA before and after the given non-coding region as context. The complete fragment of DNA including the context on both side, is then tokenized to form our input sequence of tokens. The task is to predict 919 chromatin profile including 690 transcription factors (TF) binding profiles for 160 different TFs, 125 DNase I sensitivity (DHS) profiles and 104 histone-mark (HM) profiles

subsequences, 8 subsequences out of 20). These parameters enable generating a negative set that has 32 and 40% of its sequences containing conserved portions of promoter sequences.

We prefix and append each example with [CLS] and [SEP] token respectively. The output corresponding to the [CLS] token from BIGBIRD transformer encoder is fed to a simple binary classification layer. We fine-tune the pretrained BIGBIRD from App. F.1 using hyperparameters described in Tab. 21. We note that high performance is not surprising due to the overlap in the nature of negative example generation and MLM pretraining.

### F.3 Chromatin-Profile Prediction

The first step of sequence-based algorithmic framework for predicting non-coding effects is to build a model to predict, large scale chromatic profile [109]. In this paper, we use the dataset provided in Zhou and Troyanskaya [109]<sup>7</sup>, to train BIGBIRD to predict the chromatic profile.

Each training sample consists of a 8,000-bp sequence from the human GRCh37 reference genome centered on each 200-bp bin and is paired with a label vector for 919 chromatin features. As before, we prefix and append each example with [CLS] and [SEP] token respectively. The output corresponding to the [CLS] token from BIGBIRD transformer encoder is fed to a linear layer with 919 heads. Thus we jointly predict the 919 independent binary classification problems. We fine-tune the pretrained BIGBIRD from App. F.1 using hyper-parameters described in Tab. 21. As the data is highly imbalanced data (way more negative examples than positive examples), we upweighted loss function for positive examples by factor of 8.

<sup>7.</sup> http://deepsea.princeton.edu/media/code/deepsea\_train\_bundle.v0.9.tar.gz

Parameter	Pretraining	Promoter Region	Chromatin-Profile
Block length, b	64	64	64
Global token location	ITC	ITC	ITC
# of global token, $g$	$2 \times b$	$2 \times b$	$2 \times b$
Window length, $w$	$3 \times b$	$3 \times b$	$3 \times b$
# of random token, $r$	$3 \times b$	$3 \times b$	$3 \times b$
Max. Sequence Length	4096	4096	4096
# of heads	12	12	12
# of hidden layers	12	12	12
Hidden layer size	768	768	768
Batch Size	256	256	256
Vocab Size	32000	32000	32000
Loss	$_{\rm MLM+NSP}$	BCE	919 x +ve upweighted BCE
Dropout prob	0.1	0.1	0.1
Optimizer	Adam	Adam	Adam
Learning rate	0.0001	0.0001	0.0001
# of steps	1000000	711	500000
Compute Resources	$8\times 8~\mathrm{TPUv3}$	$8 \times 8 \text{ TPUv3}$	$8 \times 8 \text{ TPUv3}$

Table 21: Table of hyperparameters for Computational biology.

We used training and testing split provided by Zhou and Troyanskaya [109] using chromosomes and strictly non-overlapping. Chromosome 8 and 9 were excluded from training to test chromatin feature prediction performances, and the rest of the autosomes were used for training and validation. 4,000 samples on chromosome 7 spanning the genomic coordinates 30,508,751–35,296,850 were used as the validation set.

As the predicted probability for each sequence in DeepSea Zhou and Troyanskaya [109] was computed as the ensemble average of the probability predictions for the forward and complementary sequence pairs, we also predict using an ensemble of two BIGBIRD model trained independently.