# **Decoder-Only Transformers**

Notes on various aspects of Decoder-Only Transformers.

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# 1 Decoder-Only Fundamentals

The Transformers architecture [1], which dominates Natural Language Processing (NLP) as of July 2023, is a relatively simple architecture. There are various flavors and variants of Transformers, but we start by describing the decoder-only versions which underly the GPT models [2–4].

The full decoder-only architecture can be seen in Fig. 1. The parameters which define the network can be found in App. A.

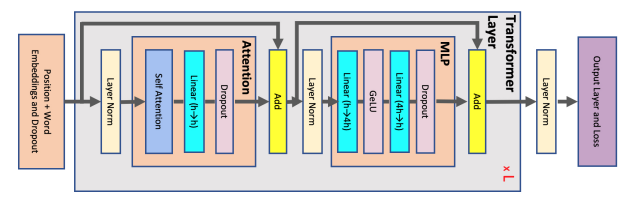


Figure 1. The full transformers architecture. Diagram taken from [5]

An outline of the mechanics:

- 1. Raw text is **tokenized** and turned into a series of integers<sup>1</sup> whose values lie in range(V), with V the vocabulary size.
- 2. The tokenized text is chunked and turned into (B, S)-shaped (batch size and sequence length, respectively) integer tensors,  $x_{bs}$ .
- 3. The **embedding layer** converts the integer tensors into continuous representations of shape (B, S, D),  $z_{bsd}$ , with D the size of the hidden dimension. **Positional encodings** have also been added to the tensor at this stage to help the architecture understand the relative ordering of the text.
- 4. The  $z_{bsd}$  tensors pass through a series of transformer blocks, each of which has two primary components:
  - (a) In the **attention** sub-block, components of  $z_{bsd}$  at different positions (s-values) interact with each other, resulting in another (B, S, D)-shaped tensor,  $z'_{bsd}$ .
  - (b) In the **MLP** block, each position in  $z'_{bsd}$  is processed independently and in parallel by a two-layer feed-forward network, resulting once more in a (B, S, D)-shaped tensor.

Importantly, there are **residual connections** around each of these<sup>2</sup> (the arrows in Fig. 1).

- 5. Finally, we convert the (B, S, D)-shaped tensors to (B, S, V)-shaped ones,  $y_{bsv}$ . This is the role of the **language model head** (which is often just the embedding layer used in an inverse manner.)
- 6. The  $y_{bsv}$  predict what the next token will be, having seen the **context** of the first s tokens in the sequence.

Each batch (the b-index) is processed independently. We omitted LayerNorm and Dropout layers above, as well as the causal mask; these will be covered below as we step through the architecture in more detail.

#### 1.1 Component Details

We break down the various components below.

#### 1.1.1 Embedding Layer and Positional Encodings

The **embedding** layer is just a simple look up table: each of the range (V) indices in the vocabulary is mapped to a D-dimensional vector via a large (V, D)-shaped table/matrix. This layer maps  $x_{bs} \longrightarrow z_{bsd}$ . In torch, this is an nn Embedding (V, D) instance.

To each item in a batch, we add identical **positional encodings** to the vectors above with the goal of adding fixed, position-dependent correlations in the sequence dimension which will hopefully

<sup>&</sup>lt;sup>1</sup>There are about 1.3 tokens per word, on average.

<sup>&</sup>lt;sup>2</sup>This gives rise to the concept of the **residual stream** which each transformer block reads from and writes back to repeatedly.

make it easier for the architecture to pick up on the relative positions of the inputs <sup>3</sup> This layer maps  $z_{bsd} \leftarrow z_{bsd} + p_{sd}$ , with  $p_{sd}$  the positional encoding tensor.

The above components require  $(V+S)D \approx VD$  parameters per layer.

#### 1.1.2 Layer Norm

The original transformers paper [1] put LayerNorm instances after the **attention** and **MLP** blocks, but now it is common [6] to put them before these blocks<sup>4</sup>.

The LayerNorm operations acts over the sequence dimension. Spelling it out, given the input tensor  $z_{bsd}$  whose mean and variance over the s-index are  $\mu_{bd}$  and  $\sigma_{bd}$ , respectively, the LayerNorm output is

$$z_{bsd} \leftarrow \left(\frac{z_{bsd} - \mu_{bd}}{\sigma_{bd}}\right) \times \gamma_d + \beta_d \equiv \text{LayerNorm }_s z_{bsd}$$
 (1.1)

where  $\gamma_d, \beta_d$  are the trainable scale and bias parameters. In torch, this is a nn.LayerNorm(D) instance.

Since there are two LayerNorm instances in each transformer block, these components require 2D parameters per layer.

#### 1.1.3 Attention

The **attention** layer is the most complex layer. It features A triplets<sup>5</sup> of weight matrices<sup>6</sup>  $Q_{df}^a, K_{df}^a, V_{df}^a$  where  $a \in \{0, \ldots, A-1\}$  and  $f \in \{0, \ldots, D/A\}$ . From these, we form three different vectors:

$$q_{bsf}^a = z_{bsd}Q_{df}^a$$
,  $k_{bsf}^a = z_{bsd}K_{df}^a$ ,  $v_{bsf}^a = z_{bsd}V_{df}^a$  (1.2)

These are the query, key, and value tensors, respectively.

Using the above tensors, we will then build up an **attention map**  $w_{bss'}^a$  which corresponds to how much attention the token at position s pays to the token at position s'. Because we have the goal of predicting the next token in the sequence, we need these weights to be causal: the final prediction  $y_{bsv}$  should only have access to information propagated from positions  $x_{bs'v}$  with  $s' \leq s$ . This corresponds to the condition that  $w_{bss'}^a = 0$  if s' > s.

These weights come from Softmax-ed attention scores, which are just a normalized dot-product over the hidden dimension:

$$w_{bss'd}^{a} = \text{Softmax}_{s'} \left( m_{ss'} + q_{bsf}^{a} k_{bs'f}^{a} / \sqrt{D/A} \right) , \text{ s.t. } \sum_{s'} w_{bdss'}^{a} = 1$$
 (1.3)

<sup>&</sup>lt;sup>3</sup>Positional encodings and the causal mask are the only components in the transformers architecture which carry weights with a dimension of size S; i.e. they are the only parts that have explicit sequence-length dependence. A related though experiment: you can convince yourself that if the inputs  $z_{bsd}$  were just random noise, the transformers architecture would not be able to predict the s-index of each such input in the absence of positional encodings.

<sup>&</sup>lt;sup>4</sup>Which makes intuitive sense for the purposes of stabilizing the matrix multiplications in the blocks

 $<sup>^{5}</sup>A$  must divide the hidden dimension D evenly.

<sup>&</sup>lt;sup>6</sup>There are also bias terms, but we will often neglect to write them explicitly or account for their (negligible) parameter count.

The tensor  $m_{ss'}$  is the causal mask which zeroes out the relevant attention map components above

$$m_{ss'} = \begin{cases} 0 & s \le s' \\ -\infty & = s > s' \end{cases}$$

The  $\sqrt{D/A}$  normalization is motivated by demanding that the variance of the Softmax argument be 1 at initialization, assuming that other components have been configured so that that the query and key components are i.i.d. from a Gaussian normal distribution <sup>7</sup>.

The weights above are then passed through a dropout layer and used to re-weigh the **value** vectors and form the tensors

$$t_{bsf}^{a} = \text{Drop}\left(w_{bdss'}^{a}\right) v_{bs'f}^{a} \tag{1.4}$$

and these A different (B, S, D/A)-shaped tensors are then concatenated along the f-direction to re-form a (B, S, D)-shaped tensor<sup>8</sup>

$$u_{bsd} = \operatorname{Concat}_{fd}^{a} \left( t_{bsf}^{a} \right) . \tag{1.5}$$

Finally, another weight matrix  $O_{d'd}$  and dropout layer transform the output once again to get the final output

$$z_{bsd} = \text{Drop} \left( u_{bsd'} O_{d'd} \right) . \tag{1.6}$$

For completeness, the entire operation in condensed notation with indices left implicit is:

$$z \leftarrow \text{Drop}\left(\text{Concat}\left(\text{Drop}\left(\text{Softmax}\left(\frac{(z \cdot Q^a) \cdot (z \cdot K^a)}{\sqrt{D/A}}\right)\right) \cdot z \cdot V^a\right) \cdot O\right)$$
 (1.7)

where all of the dot-products are over feature dimensions (those of size D or D/A). A nice<sup>9</sup>, but hopefully pedagogically useful, implementation of the attention layer is below:

```
class CausalAttention(nn.Module):
10
11
         def __init__(
12
             self,
             attn_heads=A,
13
             hidden_dim=D,
14
             block_size=K,
15
             dropout=0.1,
16
         ):
17
             super().__init__()
18
             self.head_dim, remainder = divmod(hidden_dim, attn_heads)
19
             assert not remainder, "attn_heads must divide hidden_dim evenly"
20
21
             self.Q = nn.ModuleList([nn.Linear(hidden_dim, self.head_dim) for _ in range(attn_heads)])
22
```

<sup>&</sup>lt;sup>7</sup>However, in [7] it is instead argued that no square root should be taken in order to maximize the speed of learning via SGD.

<sup>&</sup>lt;sup>8</sup>It is hard to come up with good index-notation for concatenation.

<sup>&</sup>lt;sup>9</sup>An example optimization: there is no need to form separate  $Q^a, K^a, V^a$  Linear layers, one large layer which is later chunked is more efficient

```
self.K = nn.ModuleList([nn.Linear(hidden_dim, self.head_dim) for _ in range(attn_heads)])
23
             self.V = nn.ModuleList([nn.Linear(hidden_dim, self.head_dim) for _ in range(attn_heads)])
24
             self.0 = nn.Linear(hidden_dim, hidden_dim)
25
26
             self.attn_dropout = nn.Dropout(dropout)
27
28
             self.final_dropout = nn.Dropout(dropout)
             self.register_buffer(
29
                 "causal_mask",
30
                 torch.tril(torch.ones(block_size, block_size)[None]),
31
32
33
         def get_qkv(self, inputs):
34
             queries = [q(inputs) for q in self.Q]
35
             keys = [k(inputs) for k in self.K]
36
             values = [v(inputs) for v in self.V]
37
             return queries, keys, values
38
39
         def get_attn_maps(self, queries, keys, values, seq_len):
40
41
             norm = math.sqrt(self.head_dim)
             non_causal_attn_scores = [(q @ k.transpose(-2, -1)) / norm for q, k in zip(queries, keys)]
42
             causal_attn_scores = [
43
                 a.masked_fill(self.causal_mask[:, :S, :S] == 0, float("-inf"))
44
                 for a in non_causal_attn_scores
45
46
             attn_maps = [a.softmax(dim=-1) for a in causal_attn_scores]
             return attn_maps
48
49
         def forward(self, inputs):
50
             S = inputs.shape[1]
51
             queries, keys, values = self.get_qkv(inputs)
52
             attn_maps = self.get_attn_maps(queries, keys, values, S)
53
54
             weighted_values = torch.concatenate(
55
                  [self.attn_dropout(a) @ v for a, v in zip(attn_maps, values)], dim=-1
56
             z = self.final_dropout(self.O(weighted_values))
57
```

The parameter count is dominated by the weight matrices which carry  $4D^2$  total parameters per layer.

#### 1.1.4 MLP

The feed-forward network is straightforward and corresponds to

$$z_{bsd} \leftarrow \phi \left( z_{bsd'} W_{d'e}^0 \right) W_{ed}^1 \tag{1.8}$$

where  $W^0$  and  $W^1$  are (D, FD)- and (FD, D)-shaped matrices, respectively (see App. A for notation) and  $\phi$  is a non-linearity  $^{10}$ . The implementation is straightforward:

```
8 class MLP(nn.Module):
9    def __init__(
10    self,
```

<sup>&</sup>lt;sup>10</sup>The GeLU non-linearity is common.

```
hidden_dim=D,
11
12
             expansion_factor=F,
             dropout=0.1,
13
14
15
             super().__init__()
16
             linear_1 = nn.Linear(hidden_dim, expansion_factor * hidden_dim)
             linear_2 = nn.Linear(expansion_factor * hidden_dim, hidden_dim)
17
             gelu = nn.GELU()
18
             drop = nn.Dropout(dropout)
19
             self.layers = nn.Sequential(linear_1, gelu, linear_2, drop)
20
21
         def forward(self, inputs):
22
23
             z = self.layers(inputs)
             return z
```

This bock requires  $2FD^2$  parameters per layer, only counting the contribution from weights.

#### 1.1.5 Language Model Head

The layer which converts the (B, S, D)-shaped outputs,  $z_{bsd}$ , to (B, S, V)-shaped predictions over the vocabulary,  $y_{bsv}$ , is the **Language Model Head**. It is a linear layer, whose weights are usually tied to be exactly those of the initial embedding layer of Sec. 1.1.1.

#### 1.1.6 All Together

It is then relatively straightforward to tie everything together. In code, we can first create a transformer block like

```
class TransformerBlock(nn.Module):
10
         def __init__(
11
             self,
12
             attn_heads=A,
13
             block_size=K,
14
15
             dropout=0.1,
             expansion_factor=F,
16
             hidden_dim=D,
17
             layers=L,
18
             vocab_size=V,
19
         ):
20
             super().__init__()
21
             self.attn_ln = nn.LayerNorm(hidden_dim)
22
             self.mlp_ln = nn.LayerNorm(hidden_dim)
23
             self.attn = CausalAttention(attn_heads, hidden_dim, block_size, dropout)
24
             self.mlp = MLP(hidden_dim, expansion_factor, dropout)
25
26
         def forward(self, inputs):
27
28
             z = self.attn(self.attn_ln(inputs)) + inputs
29
             z = self.mlp(self.mlp_ln(z)) + z
30
             return z
```

And then the entire architecture:

```
class DecoderOnly(nn.Module):
9
         def __init__(
10
             self,
11
             attn_heads=A,
12
             block_size=K,
13
14
             dropout=0.1,
             expansion_factor=F,
15
             hidden_dim=D,
16
             layers=L,
17
             vocab_size=V,
18
         ):
19
             super().__init__()
20
             self.embedding = nn.Embedding(vocab_size, hidden_dim)
21
             self.pos_encoding = nn.Parameter(torch.randn(1, block_size, hidden_dim))
22
             self.drop = nn.Dropout(dropout)
23
             self.trans_blocks = nn.ModuleList(
24
                  Γ
25
26
                      TransformerBlock(
27
                          attn_heads,
28
                          block_size,
                          dropout,
29
                          expansion_factor,
30
                          hidden_dim,
31
                          layers,
32
                          vocab_size,
33
                      )
34
                      for _ in range(layers)
35
                 ]
36
             )
37
             self.final_ln = nn.LayerNorm(hidden_dim)
38
             self.lm_head = nn.Linear(hidden_dim, vocab_size, bias=False)
39
40
             self.lm_head.weight = self.embedding.weight # Weight tying.
41
         def forward(self, inputs):
42
             S = inputs.shape[1]
43
             z = self.embedding(inputs) + self.pos_encoding[:, :S]
44
             z = self.drop(z)
45
46
             for block in self.trans_blocks:
                 z = block(z)
47
             z = self.final_ln(z)
48
             z = self.lm_head(z)
49
             return z
50
51
```

#### 1.2 Memory

Understanding the memory cost of

## A Conventions and Notation

We loosely follow the conventions of [5] and denote the main Transformers parameters by:

• B: microbatch size

- K: the block size (maximum sequence length  $^{11}$ )
- S: input sequence length
- V: vocabulary size
- D: the hidden dimension size
- L: number of transformer layers
- A: number of attention heads
- P: pipeline parallel size
- T: tensor parallel size
- F: expansion factor for MLP layer (usually F = 4)

Where it makes sense, we try to use the lower-case versions of these characters to denote the corresponding indices on various tensors. For instance, an input tensor with the above batch size, sequence length, and vocabulary size would be written as  $x_{bsv}$ , with  $b \in \{0, ..., B-1\}$ ,  $s \in \{0, ..., S-1\}$ , and  $v \in \{0, ..., V-1\}$  in math notation, or as x[b, s, v] in code. Typical transformers belong to the regime

$$V \gg D, S \gg L, A \gg P, T$$
 (A.1)

As indicated above, we use zero-indexing. We also use python code throughout  $^{12}$  and write all ML code using standard torch syntax. To avoid needing to come up with new symbols in math expressions we will often use expressions like  $x \leftarrow f(x)$  to refer to performing a computation on some argument (x) and assigning the result right back to the variable x again.

Physicists often joke (half-seriously) that Einstein's greatest contribution to physics was his summation notation in which index-sums are implied by the presence of repeated indices and summation symbols are entirely ommitted. For instance, the dot product between two vectors would be written as

$$\vec{x} \cdot \vec{y} = \sum_{i} x_i y_i \equiv x_i y_i \tag{A.2}$$

We use similar notation which is further adapted to the common element-wise deep-learning operations. The general rule is that if a repeated index appears on one side of an equation, but not the other, then a sum is implied, but if the same index appears on both sides, then it's an element-wise operation. The Hadamard-product between two matrices A and B is just

$$C_{ij} = A_{ij}B_{ij} . (A.3)$$

We also put explicit indices on operators such as Softmax to help clarify the relevant dimension, e.g. we would write the softmax operation over the b-ndex of some batched tensor  $x_{bvd...}$  as

$$s_{bvd\dots} = \frac{e^{x_{bvd\dots}}}{\sum_{v=0}^{v=V-1} e^{x_{bvd\dots}}} \equiv \text{Softmax}_{v} x_{bvd\dots}, \qquad (A.4)$$

indicating that the sum over the singled-out v-index is gives unity.

<sup>&</sup>lt;sup>11</sup>In the absence of methods such as ALiBi [8] can be used to extend the sequence length at inference time.

<sup>&</sup>lt;sup>12</sup>Written in a style conducive to latex, e.g. no type-hints and pegagogy prioritized.

### B TODO

- Tokenizers
- Generation
- Activations
- Flash attention
- BERT family
- Residual stream
- Scaling laws

#### References

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