



# Eukaryotic and prokaryotic gene structure

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## Abstract

Genes consist of multiple sequence elements that together encode the functional product and regulate its expression. Despite their fundamental importance, there are few freely available diagrams of gene structure. Presented here are two figures that summarise the different structures found in eukaryotic and prokaryotic genes. Common gene structural elements are colour-coded by their function in regulation, transcription, or translation.

## Introduction

### Gene structure

Genes contain the information necessary for living cells to survive and reproduce.<sup>[1][2]</sup> In most organisms, genes are made of DNA, where the particular DNA sequence determines the function of the gene. A gene is transcribed (copied) from DNA into RNA, which can either be non-coding (ncRNA) with a direct function, or an intermediate messenger (mRNA) that is then translated into protein. Each of these steps is controlled by specific sequence elements, or regions, within the gene. Every gene, therefore, requires multiple sequence elements to be functional.<sup>[2]</sup> This includes the sequence that actually encodes the functional protein or ncRNA, as well as multiple regulatory sequence regions. These regions may be as short as a few base pairs, up to many thousands of base pairs long.

Much of gene structure is broadly similar between eukaryotes and prokaryotes. These common elements largely result from the shared ancestry of cellular life in organisms over 2 billion years ago.<sup>[3]</sup> Key differences in gene structure between eukaryotes and prokaryotes reflect their divergent transcription and translation machinery.<sup>[4][5]</sup> Understanding gene structure is the foundation of understanding gene annotation, expression, and function.<sup>[6]</sup>

### Previous images

Access to freely available diagrams is important for scientists, medical professions and the general public.<sup>[7][8]</sup> However, current open-access gene structure figures are limited in their scope (Supplementary figures 1-4), typically showing one or a few aspects (e.g. exon splicing, or promoter regions). Information-rich diagrams that use a consistent colour scheme and layout should help the comprehension of complex concepts.<sup>[9]</sup>

This work provides two diagrams that summarise the complex structure and terminology of genes. Common elements of gene structure are presented in a consistent layout and format to highlight the relationships between components. Key differences between eukaryotes and prokaryotes are indicated.

## Results

### Common gene structure features

The structures of both eukaryotic and prokaryotic genes involve several nested sequence elements. Each element has a specific function in the multi-step process of gene expression. The sequences and lengths of these elements vary, but the same general functions are present in most genes.<sup>[2]</sup> Although DNA is a double-stranded molecule, typically only one of the strands encodes information that the RNA polymerase reads to produce protein-coding mRNA or non-coding RNA. This 'sense' or 'coding' strand, runs in the 5' to 3' direction where the numbers refer to the carbon atoms of the backbone's ribose sugar. The open reading frame (ORF) of a gene is therefore usually represented as an

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Received 14-11-2016; accepted 17-01-2017



arrow indicating the direction in which the sense strand is read.<sup>[10]</sup>

**Regulatory sequences** are located at the extremities of genes. These sequence regions can either be next to the transcribed region (the **promoter**) or separated by many kilobases (**enhancers** and **silencers**).<sup>[11]</sup> The promoter is located at the 5' end of the gene and is composed of a core promoter sequence and a proximal promoter sequence. The core promoter marks the start site for transcription by binding RNA polymerase and other proteins necessary for copying DNA to RNA. The proximal promoter region binds **transcription factors** that modify the affinity of the core promoter for RNA polymerase.<sup>[12][13]</sup> Genes may be regulated by multiple enhancer and silencer sequences that further modify the activity of promoters by binding **activator** or **repressor** proteins.<sup>[14][15]</sup> Enhancers and silencers may be distantly located from the gene, many thousands of base pairs away. The binding of different transcription factors, therefore, regulates the rate of transcription initiation at different times and in different cells.<sup>[16]</sup>

Regulatory elements can overlap one another, with a section of DNA able to interact with many competing activators and repressors as well as RNA polymerase. For example, some repressor proteins can bind to the core promoter to prevent polymerase binding.<sup>[17]</sup> For genes with multiple regulatory sequences, the rate of transcription is the product of all of the elements combined.<sup>[18]</sup> Binding of activators and repressors to multiple regulatory sequences has a cooperative effect on transcription initiation.<sup>[19]</sup>

Although all organisms use both transcriptional activators and repressors, eukaryotic genes are said to be 'default off', whereas prokaryotic genes are 'default on'.<sup>[5]</sup> The core promoter of eukaryotic genes typically requires additional activation by promoter elements for expression to occur. The core promoter of prokaryotic genes, conversely, is sufficient for strong expression and is regulated by repressors.<sup>[5]</sup>

An additional layer of regulation occurs for protein coding genes after the mRNA has been processed to prepare it for translation to protein. Only the region between the **start** and **stop** codons encodes the final protein product. The flanking **untranslated regions** (UTRs) contain further regulatory sequences.<sup>[20]</sup> The **3' UTR** contains a **terminator** sequence, which marks the endpoint for transcription and releases the RNA polymerase.<sup>[21]</sup> The **5' UTR** binds the **ribosome**, which translates the **protein-coding region** into a string of **amino acids** that **fold** to form the final protein product. In the case of genes for non-coding RNAs the RNA is not translated but instead **folds** to be directly functional.<sup>[22][23]</sup>

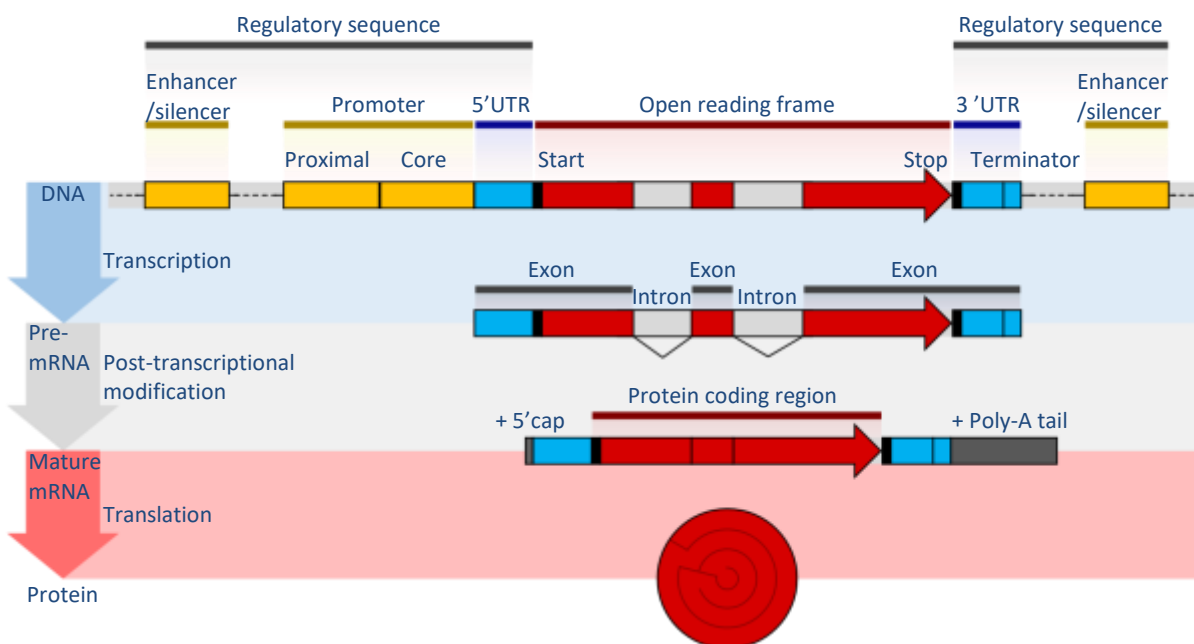
## Eukaryotes

The structure of eukaryotic genes includes features not found in prokaryotes (Figure 1). Most of these relate to **post-transcriptional modification** of **pre-mRNAs** to produce **mature mRNA** ready for translation into protein. Eukaryotic genes typically have more regulatory elements to control gene expression compared to prokaryotes.<sup>[5]</sup> This is particularly true in **multicellular** eukaryotes, humans for example, where gene expression varies widely among different **tissues**.<sup>[24]</sup>

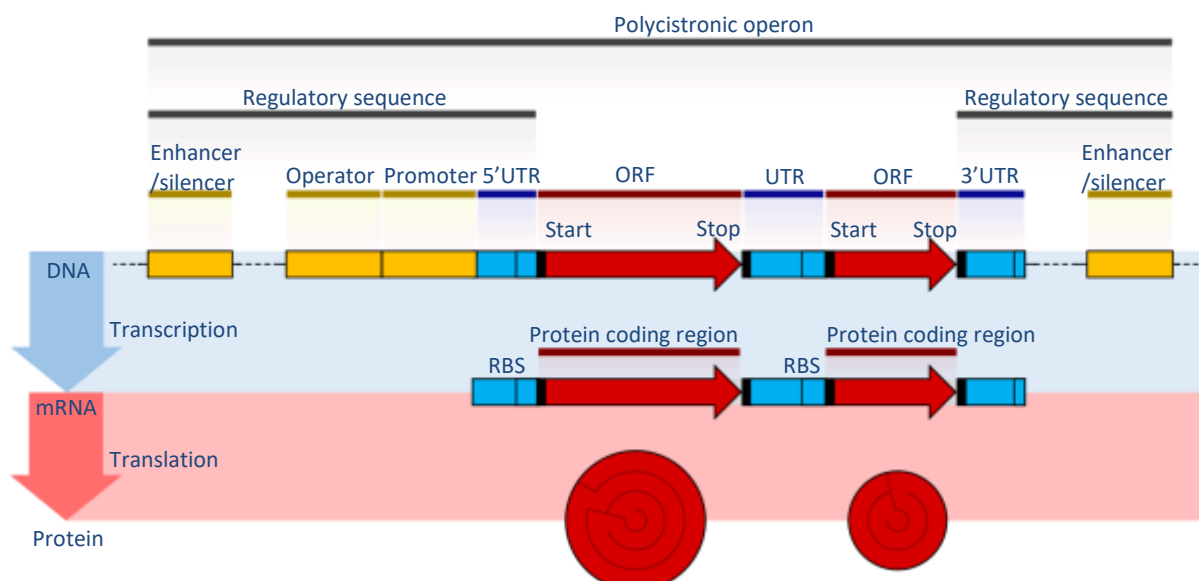
A key feature of the structure of eukaryotic genes is that their transcripts are typically subdivided into **exon** and **intron** regions. Exon regions are retained in the final **mature mRNA** molecule, while intron regions are **spliced out** (excised) during post-transcriptional processing.<sup>[25]</sup> Indeed, the intron regions of a gene can be considerably longer than the exon regions. Once spliced together, the exons form a single continuous protein-coding regions, and the splice boundaries are not detectable. Eukaryotic post-transcriptional processing also adds a **5' cap** to the start of the mRNA and a **poly-adenosine tail** to the end of the mRNA. These additions stabilise the mRNA and direct its **transport** from the **nucleus** to the **cytoplasm**, although neither of these features are directly encoded in the structure of a gene.<sup>[20]</sup>

## Prokaryotes

The overall organisation of prokaryotic genes is markedly different from that of the eukaryotes (Figure 2). The most obvious difference is that prokaryotic ORFs are often grouped into a **polycistronic operon** under the control of a shared set of regulatory sequences. These ORFs are all transcribed onto the same mRNA and so are co-regulated and often serve related functions.<sup>[26][27]</sup> Each ORF typically has its own **ribosome binding site** (RBS) so that ribosomes simultaneously translate ORFs on the same mRNA. Some operons also display translational coupling, where the translation rates of multiple ORFs within an operon are linked.<sup>[28]</sup> This can occur when the ribosome remains attached at the end of an ORF and simply translocates along to the next without the need for a new RBS.<sup>[29]</sup> Translational coupling is also observed when translation of an ORF affects the accessibility of the next RBS through changes in RNA secondary structure.<sup>[30]</sup> Having multiple ORFs on a single mRNA is only possible in prokaryotes because their transcription and translation take place at the same time and in the same subcellular location.<sup>[26][31]</sup>



**Figure 1 |** The structure of a eukaryotic protein-coding gene. Regulatory sequence controls when and where expression occurs for the protein coding region (red). Promoter and enhancer regions (yellow) regulate the transcription of the gene into a pre-mRNA which is modified to remove introns (light grey) and add a 5' cap and poly-A tail (dark grey). The mRNA 5' and 3' untranslated regions (blue) regulate translation into the final protein product.



**Figure 2 |** The structure of a prokaryotic operon of protein-coding genes. Regulatory sequence controls when expression occurs for the multiple protein coding regions (red). Promoter, operator and enhancer regions (yellow) regulate the transcription of the gene into an mRNA. The mRNA untranslated regions (blue) regulate translation into the final protein products.



The **operator** sequence next to the promoter is the main regulatory element in prokaryotes. Repressor proteins bind to the operator sequence physically obstructs the RNA polymerase enzyme, preventing transcription.<sup>[32][33]</sup> **Riboswitches** are another important regulatory sequence commonly present in prokaryotic UTRs. These sequences switch between alternative secondary structures in the RNA depending on the concentration of key **metabolites**. The secondary structures then either block or reveal important sequence regions such as RBSs. Introns are extremely rare in prokaryotes and therefore do not play a significant role in prokaryotic gene regulation.<sup>[34]</sup>

## Discussion

### Strengths and limitations

The structures of genes are intimately linked to their functions. Genes are often thought of as only the protein-coding regions of DNA. However, coding sequences are only a minority component of the overall gene. Multiple untranscribed and untranslated DNA regions are necessary for proper gene function in all organisms. All regions of gene structure are important for determining phenotype, and so mutations in any of these sections have the potential to alter function.<sup>[35][36]</sup>

These two figures aim to present the key features of average genes. As such, they inevitably miss many interesting features, such as **overlapping genes**,<sup>[37]</sup> **alternative splicing**,<sup>[38]</sup> **trans-splicing**,<sup>[39]</sup> or the **subcellular locations** of events.<sup>[6]</sup> All the functions performed by the sequences shown are mediated through their interactions with pre-existing **proteins** and other **trans-acting elements**, which have been omitted for clarity. DNA is similarly stored in complex with **histone** proteins. Changes in this **quaternary structure** by **epigenetic** mechanisms such as **histone modification** or **DNA methylation** also affect how accessible genes are to the transcription machinery.<sup>[40]</sup> Despite these concessions, the diagrams provide an accurate overview of the important common aspects of gene structure necessary for understanding genetics.

### Future extensions

These diagrams have been designed to be extended to other systems in a consistent visual format. For example, in addition to cellular gene structure, viruses have highly variable and complex genes with features dependent on their specific host. A diagram of gene features found in viruses would be of benefit to the community and could feature either model viruses, such as

bacteriophage, or medically important examples, such as influenza or HIV.

RNA transcripts contain post-transcriptional modifications that are beyond the scope of the figures presented here. A figure outlining the RNA modifications that control transcript stability, transport, and translation would be a useful extension to these diagrams and extend the series from DNA into RNA structure. The transcriptional complex assembly and the spliceosomal complex assembly are two additional biological phenomena that would be well suited to this series. The protein components required and the order of assembly is well described yet the number of diagrams on Wikipedia is limited at this stage. Future works should extend the series by adding new concepts within the same unified format.

### Conclusion

While these diagrams cover well established biological processes, there was an unmet need for logical diagrams with the clarity for readers to learn the basic concepts of gene structure. This work improves over existing representations by linking unifying concepts between eukaryotes and prokaryotes and explaining the concepts in a clear layout.

### Availability

The diagrams are freely available in the following formats:

	Eukaryote		Prokaryote	
	Hyper-linked	Non-hyper-linked	Hyper-linked	Non-hyper-linked
svg	<a href="#">link</a>	<a href="#">link</a>	<a href="#">link</a>	<a href="#">link</a>
Wikimarkup	<a href="#">link</a>	-	<a href="#">link</a>	-
png	-	<a href="#">link</a>	-	<a href="#">link</a>

Unannotated versions for use in translations are available [here](#) and [here](#).

### Acknowledgements

The authors would like to thank [Adrian J. Hunter](#), [Boghog](#), and [Opabinia regalis](#) for their excellent recommendations during the [Good Article review](#) of Wikipedia's [Gene](#) page.

### Declarations

Conflict of Interest: TS is a member of the Editorial Board of *WikiJ.Med.* and has absented himself from all discussion and processing of this manuscript. Otherwise, we have no conflicts of interest.



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