Chromatin properties of temperature-adapted eukaryotes

David Albir Haba
Title: Chromatin properties of temperature-adapted eukaryotes

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Abstract:

Chromatin packaging is linked with the genome sequence, molecular mechanisms and biological properties of cells. We hypothesize that the histone proteins packaging cold-living eukaryotes (psychrophiles) have enhanced flexibility over mesophiles such as commonly studied animals and fungi, whereas the histones of warm adapted eukaryotes (thermophiles) have enhanced stability compared to mesophiles.

In this project we will use synthetic genes to recombinantly express and purify histones with sequence of the Antarctic fish Notothenia coriiceps living below 4ºC and the fungus Chaetomium thermophilum with optimum living temperature of 60ºC.

We attempted to assemble these into histone octamers and nucleosomes to enable comparison of their properties with human equivalents.

Key words: Histones, Temperature-adapted, Chromatin, Eukaryotes, Fungi, Bacteria, Thermophiles, Mesophiles.
Titúlo: Propiedades de la cromatina en eucariotas adaptados a la temperatura

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Resumen:

La compactación de la cromatina está relacionada con la secuencia genómica, los mecanismos moleculares y las propiedades biológicas de las células. Se hipotetiza que el compactamiento de las proteínas histonas de los animales eucariotas adaptados al frio (psicrófilos) han ganado en flexibilidad frente a los mesófilos. En cambio las histonas de los eucariotas adaptados a altas temperaturas (termófilos) como los hongos han ganado en estabilidad en comparación frente a los mesófilos.

En este proyecto genes recombinantes serán usados para expresar y purificar las histonas del pez de la Antartida Notothenia coriiceps que vive por debajo de los 4ºC y el hongo Chaetomium thermophilum con una temperatura óptima de desarrollo de 60ºC.

Se ensamblaran estas histonas en octámeros y nucleosomas para permitir la comparación de sus propiedades con sus equivalentes humanos.

Palabras clave:

Histonas, Adaptación-temperatura, Cromatina, Eucariotas, Hongos, Bacterias, Termófilos, Psicrófilos, Mesófilos
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INDEX

1. INTRODUCTION ............................................................................................................. 3
   1.1. Characteristics and thermal adaptation of eukaryotes ............................................. 3
       1.1.1. Psychrophiles ................................................................................................. 4
       1.1.2. Thermophiles ............................................................................................... 5
   1.2. Functions of chromatin in eukaryotes ..................................................................... 6
       1.2.1. Structure of chromatin ................................................................................ 7
       1.2.2. Epigenetics .................................................................................................. 8
   1.3. Histones and nucleosomes ..................................................................................... 8
       1.3.1. Histone proteins ......................................................................................... 8
       1.3.2. Histone fold dimers and octamers ............................................................... 9
       1.3.3. Nucleosomes ............................................................................................. 9
   1.4. Aims and objectives .............................................................................................. 10

2. MATERIALS AND METHODS ..................................................................................... 11
   2.1. Materials .............................................................................................................. 11
   2.2. Methods ............................................................................................................... 13
       2.2.1. DNA methods ............................................................................................ 13
       2.2.2. Protein methods ....................................................................................... 14
       2.2.3. Nucleosome methods ............................................................................... 17

3. RESULTS .................................................................................................................... 18
   3.1. Establishing novel histone expression .................................................................. 18
       3.1.1. De novo sequence optimization ................................................................ 18
       3.1.2. Plasmid construction ................................................................................. 18
       3.1.3. N. coriiceps H3 mutagenesis ................................................................. 20
   3.2. Optimization of protein expression ..................................................................... 21
       3.2.1. Expression media trial .............................................................................. 21
       3.2.2. Expression strain trial ............................................................................. 21
       3.2.3. Variation in expression between inoculums ............................................. 23
       3.2.4. Optimization of C. thermophilum H4 ribosome binding site ................ 23
   3.3. Preparation of histones ....................................................................................... 25
       3.3.1. Large scale expression .............................................................................. 25
       3.3.2. Purification by chromatography ............................................................... 25
       3.3.3. Purification by gel filtration chromatography ........................................ 26
   3.4. Assembly of octamers refolding from histones .................................................. 27
3.4.1. \textit{N. coriiceps} octamers ................................................................. 28
3.4.2. \textit{C. thermophilum} histones do not refold into octamers ......................... 29
3.5. Nucleosome assembly ............................................................................ 29
3.5.1. \textit{N. coriiceps} nucleosomes in vitro by salt dialysis .............................. 30
4. DISCUSSION .......................................................................................... 31
4.1. Histone selection and gene design ............................................................ 31
4.2. Protein expression and purification .......................................................... 31
4.3. Octamers refolding and nucleosome ....................................................... 32
5. CONCLUSIONS ...................................................................................... 33
6. APPENDIX ............................................................................................. 34
6.1. Histone Genes sequences ...................................................................... 34
6.2. Histone Protein sequences .................................................................... 36
6.3. Histone proteins alignments .................................................................. 37

LIST OF FIGURES AND TABLES

\textbf{Figure 1.} Stability curve for a human protein
\textbf{Figure 2.} \textit{C. thermophilum} protein Arx1
\textbf{Figure 3.} Nucleosome compaction of DNA into
\textbf{Figure 4.} Dimers conformation.
\textbf{Figure 5.} Nucleosome dialysis
\textbf{Figure 6.} Structure of the plasmid with the histone gene CtH2AX
\textbf{Figure 7.} \textit{N. coriiceps} H3 mutagenesis sequences.
\textbf{Figure 8.} Test expression media optimization.
\textbf{Figure 9.} Test expression and media optimization.
\textbf{Figure 10.} Expression trial.
\textbf{Figure 11.} Plasmid structure marking the region where the insertion of the gap and primers Design for \textit{C. thermophilum} H4 mutagenesis
\textbf{Figure 12.} SDS-Page Gel with the mutagenesis of \textit{C. thermophilum} H4.
\textbf{Figure 13.} Anion exchange chromatography for NcH3
\textbf{Figure 14.} Process column filtration eliminate DNA.
\textbf{Figure 15.} \textit{N. coriiceps} histones refolded in octamers
\textbf{Figure 16.} \textit{C. thermophilum} histones do not refolded in octamers.
\textbf{Figure 17.} \textit{N. coriiceps} Nucleosomes and tetrasomes on 54A54 DNA sequence

\textbf{Table 1.} Solutions and Buffers
\textbf{Table 2.} PCR conditions for KOD hot start polymerase.
\textbf{Table 3.} PCR program using KOD hot start polymerase.
\textbf{Table 4.} Description of the vector with the correspondent histone gene
\textbf{Table 5.} Sumary table of origin, state, properties and production of each histone.
1. INTRODUCTION

1.1. Characteristics and thermal adaptation of eukaryotes

Eukaryote is an ancient Greek word which means ‘true nucleus’ (sic). The Eukaryote domain of life comprises cells with a compact nucleus and defined compartments separated by membranes, which are thought to be invaginations of the external plasma membrane. The nuclear envelope delimits the nucleus, which is the largest organelle in the cell and contains the genetic code of the cell in the form of chromosomes (Pollard et al., 2007). This specific compaction of DNA is distinct in the Eukaryote domain compared to the other two prokaryotic domains, Bacteria and Archaea.

The Eukaryote domain includes a great variety of organisms apparently very different from each other, that form five supergroups: Amoeboza, Excavata, Archaeplastida, Stremnopiles, Alveolata and Rhizaria (SAR) and Opisthokonta (Adl et al., 2012). These supergroups in turn, comprise kingdoms including Fungi (Chaetomium thermophilum) and Metazoan/Animata (Homo sapiens, Xenopus laevis, Caenorhabditis elegans and Notothenia coriceps) among others.

Eukaryotes and other organisms have struggled against their environment, so under Darwin’s theory they adapt, change and evolve. Temperature is one environmental variable for which adaptation has been necessary, and the impact of the temperature in Eukaryotes is well known (Krenek et al., 2012).

At the molecular level, this adaptation depends on the behavior of proteins and other biomolecules. The stability curve of a protein is defined as the plot of the free energy of unfolding as a function of temperature. Human proteins are in their most stable conformation around 25°C (Figure 1).
Figure 1. Stability curve for a human protein. The curve is obtained from temperature variation of enthalpy and entropy of unfolding (Becktel and Schellman, 1987).

A mesophile organism has an optimal living temperature between 15º and 35º C. Other organisms are specialized to survive at extreme temperatures and their adaptation is interesting. In this project we have investigated how Eukaryotic cells have solved the problem of DNA compactation in variable thermal circumstances.

1.1.1. Psychrophiles

The word psychrophiles refers to the organisms able to live at temperatures below 5ºC. However, their optimal temperatures for development can be between 12º and 15ºC, and their upper temperature tolerance can be between 15º and 20ºC.

In a 1928 expedition in Norway, Ditlef Rustad captured a translucid fish in the south shore of the island of Bouvetøya, and gave it the name of "Farveløst Blood" (‘Colorless blood’) in his diary (Rustad, 1930). Many years later it was realized that these ice-fish are the only vertebrate animals without myoglobin or hemoglobin in their blood. In these cold temperatures oxygen dissolves to a higher concentration than in mesophile temperatures so can dissolve directly into plasma, which is less viscous because it does not require erythrocytes. The absence of oxygen carrying proteins enabled the fish to develop much larger hearts, denser capillary beds, higher volumes of blood and larger blood vessels where plasma can circulate faster with lower energy expenditure.

Another adaptation of ice fish is the appearance of antifreeze proteins, which derive from an ancestral trypsinogen gene that was duplicated in an example of neofunctionalisation (Beers et al., 2015). One copy maintained the original role as protease, but the other accumulated mutations that eventually provided the antifreeze function. These proteins bind to ice crystals and prevent their growth that results in cell damage. As shown by genomic and proteomic
studies, cold-shock proteins are highly expressed in ice fish and can have crucial roles in protein folding, control of nucleic-acid secondary structure, and transcription and translation (Shin et al., 2014).

Proteins are the main targets of these adaptations as they control the equilibrium between substrates and products, influx of nutrients, outflow of waste products, macromolecular assemblies, nucleic-acid dynamics and appropriate folding. Their adaptation seems to rely on a higher flexibility of key parts of the molecular structure or of whole folded domains through a decreased stability that partly compensates for the freezing effect of low temperatures on the three-dimensional structure. The Heat Shock Response (HSR) is a mechanism of defense against thermal stress from elevated temperatures. Curiously, HSR proteins seems to be constitutively expressed in cold adapted organisms, presumably to mitigate cold denaturation of proteins (Shin et al., 2014).

Global warming does not bode well for Antarctic fish, and even less for ice-fish which are more sensitive to temperature changes than red-blooded fish. In addition, an increase in temperature leads to an increase in the acidity of the ocean and, as a result, an imbalance in ecosystems and food webs.

Despite their adaptations to the cold environment, genome-wide studies have only recently been performed due to the lack of a sequenced genome (Shin et al., 2014). This revealed that *N. coriiceps* has bigger genome, higher dN/dS ratio comparing with the other species that belong to the same family but not exposed to this environment and multiple post-translational modifications (Shin et al., 2014). The Antarctic bullhead Notother, *N. coriiceps*, is therefore an endemic teleost ice fish which can be a good model to understand and represent the cold evolution adaptations to the genome.

1.1.2. Thermophiles
Thermophile is a term applied to the organisms that can grow in extreme conditions of high temperatures above 50ºC. Such organisms have been widely used in biotechnology because of the inherent property that their proteins are resistant to high temperatures. *Chaetomium thermophilum* is a thermophilic filamentous fungus whose genome has been published recently (Kellner et al., 2016). It grows on dung or compost and has an optimum temperature of 60ºC, which is one of the highest for Eukaryotes. In contrast, *Saccharomyces cerevisiae*, a model yeast, has 73% protein homology to this fungus but has a temperature optimum of only 30ºC (van den Brink et al. 2015).

Proteins for thermophilic organisms are not only stable to unfolding at higher temperatures, but also generally more stable than their mesophilic relatives. Alongside this high stability and
resistance of denaturation of proteins, thermophiles such *C. thermophilum* show codon usage bias and reduced genome size with fewer protein coding genes than mesophiles of the same family (*C. thermophilum* 7267, *T. terrestris* 9813; *T. heterothallica* 9110). Shorter introns and intergenic regions and less transposable elements may indicate that transposition is unfavorable at higher temperatures (van den Brink et al. 2015).

On the other hand, thermophile proteins show a high concentration of isoleucine, tryptophan and tyrosine. There is also an overrepresentation of cysteine for catalytic residues, disulfide bridges and metal binding contribute to protein thermal stability, and proline which makes the structure of the proteins more rigid and less unfoldable.

**Figure 2.** *Ct. thermophilum* protein Arx1. This specific protein of *C. thermophilum*, which is homologous to yeast pre-ribosomal export factor Arx1, unfolds around 62°C. (Van Noort et al. 2013)

By solving the 3D structure of the *C. thermophilum* protein Arx1 (Figure 2), three types of adaptive mutations were uncovered: 1) the loop rigidity thanks to the increased proline frequency; 2) an increased protein core hydrophobicity, and 3) an increase in electrostatic interactions that stabilize neighboring secondary structure elements (van Noort et al. 2013).

This makes *C. thermophilum* an interesting model for adaptation because its genome and proteome are very well characterized and provide a basis to interpret how histones have evolved in order to perform the nuclear compaction of the DNA.

### 1.2. Functions of chromatin in eukaryotes

The main function of the chromatin is to package the genetic material in the eukaryotic cell. DNA is packaged in a series of repeating nucleosome units in chromosomes, and this structure is predictable and highly conserved. The conformation of the nucleosomes provides a balance
of static and dynamic organization which is essential for chromatin function. The other functions of the chromatin apart from packaging the DNA are to organize and control the expression of the proteins, DNA replication and DNA repair (Figure 3).

**Figure 3.** Nucleosome compaction of DNA into chromosomes (Pierce 2003).

**1.2.1. Structure of chromatin**

To perform special cell functions and respond to a changing environment, the way that the genome is accessible to the transcriptome and regulation complex machinery is dynamic and coordinated. Chromatin serves as platform for all nuclear processes from gene expression to DNA replication and DNA damage repair. This structure changes in part thanks to remodeling complexes that control nucleosome positioning. The profile of nucleosomes varies throughout
the cell-cycle involving histone post-translation modifications and incorporation of histone variants via chromatin remodeling complexes during gene regulation (Bruno et al., 2003).

1.2.2. Epigenetics
A variety of post-translational modifications including acetylation, phosphorylation, methylation, ubiquitination and ADP-ribosylation occur on the tail regions of histones. These modifications led to the ‘histone code hypothesis’ whereby marks in some histones tails encode changes in the chromatin-templated processes. This histone code seems to be, at least in some part, heritable or epigenetic.

Some studies show that the exposure to high temperatures for one generation in C. elegans changes the expression of temperature-induced proteins until at least the 14th generation. This change is related with the altered trimethylation of histone H3 lysine 9 (H3K9me3) before the onset of zygotic transcription. When the worms were at 20ºC, they produced a small quantity of a fluorescent protein. However, when the temperature of the habitat was raised to 25ºC, the amount of fluorescent protein produced increased. Afterwards, even after return to the lower the temperature of 20ºC, the fluorescent protein production was maintained. Most surprising was that this kind of memory of the warm period was not only maintained in individuals who experimented it but also in the children and grandchildren of these single worms maintained only at 20ºC. These later generations continued to show the fluorescence that indicated the experience of their parents and grandparents to heat. The effect lasted up to seven generations and, for a generator of five generations at 25ºC, the fluorescence was maintained up to 14 generations.

One of the explanations for this phenomenon is that because the generations of these worms are so short and the environment could change more slowly, that adaptation for long-lived epigenetic inheritance may be useful to them (Klosin et al., 2017).

1.3. Histones and nucleosomes

1.3.1. Histone proteins
Histones are small proteins of 10-15 kDa that are highly conserved in evolution. There are 5 types of histones, H2A, H2B, H3, H4 and H1. H2A has a variant, H2AX, found in almost all eukaryotes that makes up a significant but variable proportion of bulk chromatin and plays a role in DNA damage and repair. H2A and H2B form a histone fold dimer, and H3 and H4 form another one. Nucleosomes involve 2 each of the 4 core histones H2A, H2B, H3 and H4 and one molecule of H1 for every 200 bp of DNA. The nucleosome core particle (NCP) structures have been solved using histones from Xenopus and Human and other organisms before (Patwal, 2014). NCPs are connected to each other by a DNA linker of 20-100 bp, that can be associated
with a molecule of H1. It has been extremely difficult to understand the principles of binding histones to DNA because wrapped DNA behaves as an irregular and cooperative superhelix.

### 1.3.2. Histone fold dimers and octamers

The histone octamer in an NCP is an arrangement of (H3-H4), tetramer at the center and H2A-H2B dimers on either side of the tetramer. These histones are associated as heterodimers by specific hydrophobic interactions in anti-parallel pairs of H2A-H2B and H3-H4 (Figure 4).

![Histone fold dimers conformation](image)

**Figure 4. Histone fold dimers conformation.** The histones adopt this conformation this order to perform an octamer: (H2A–H2B:H4–H3:H3–H4:H2B–H2A) (Patwal, 2016)

The octamer conformation provides a characteristic spiral of histone fold dimers in a specific way that makes possible the histone–DNA interactions which is crucial for nucleosome positioning. The histone fold motif comprises a long central α2 helix flanked by shorter α1 and α3 helices. Folding forces based mainly on hydrophobic interactions allow antiparallel pairs H2A–H2B and H3–H4 to interact along the α2 helix, with the α1 and α3 helices folded back across the long α2 helix (Dechassa and Luger, 2012).

The octamer conformation is the platform enabling DNA contacts. Two types of DNA interaction are generated in the surfaces of α1 and α3 helices. The first is formed by the combination of pairs of loops connecting alpha helices known as L1L2 motifs because the pairs involve the loops between α1 and α2 (L1), and α2 and α3 (L2). The second type of interaction surface is formed by paired N-terminal ends of α1 helices, and is known as the α1α1 motif (Flaus, 2011).

### 1.3.3. Nucleosomes

The nucleosome is defined as the repeating unit of chromatin. The NCP is formed by a structured octamer of the core histones H2A, H2B, H3 and H4 wrapped with 147 bp DNA (Oudet et al., 1975). Nucleosomes are arranged into regularly spaced array, with the linker region between nucleosomes. Nucleosome positioning influences gene expression, and has
implications for chromatin packaging, sequence evolution, and the evolution of gene expression. The three principles of nucleosome dynamics and positioning are (i) DNA sequence accessibility depends on its position within the nucleosome structure, (ii) DNA sequences can influence their location in the nucleosome and (iii) DNA and the histone octamer core together control the potential for structural changes in the nucleosome. The propensity for nucleosomes to resist or undergo structural changes is referred to as nucleosome stability (Flaus, 2011).

1.4. Aims and objectives

Different types of genomes need different properties of chromatin compaction, so variations in histones are expected. However, histones and nucleosomes appear to be highly conserved in Eukaryotic evolution. Significant but small changes in histone sequences are likely to affect the compaction of the DNA and genome, especially in divergent eukaryotes such as those exposed to different thermal adaptations. We set out to compare the histones of the thermophile *C. thermophilum* and the psychrophile *N. coriiceps* with those from more well studied mesophile animals like *H. sapiens* and *X. laevis*.

The aim of this project is uncover the nature of histones variations that enable adaptation to extreme temperatures. To accomplish this, our first objective was to express all four core histones for each organism recombinantly in *E. coli* and then to purify the recombinant histones and assemble them into nucleosomes.
# 2. MATERIALS AND METHODS

## 2.1. Materials

<table>
<thead>
<tr>
<th>Buffers and solutions</th>
<th>Composition</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ampicillin</td>
<td>Stock Solution: 50 mg/ml in ddH2O. Used at 50 ug/ml</td>
<td>Selection for antibiotic resistance</td>
</tr>
<tr>
<td>Binding Buffer</td>
<td>100mM Tris-HCl pH 7.5, 500 mM NaCl and 0.1 mM EDTA</td>
<td>For binding proteins for column filtration</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>stock solution: 34 mg/ml in absolute Ethanol. Used at 34 ug/ml</td>
<td>Selection for antibiotic resistance</td>
</tr>
<tr>
<td>Coomassie Stain</td>
<td>50% v/v Methanol, 10% v/v Acetic Acid, 40% v/v deionized water, 0.1% 2wv Brilliant Blue G, 0.01% w/v Brilliant Blue R</td>
<td>Protein stain in SDS-PAGE</td>
</tr>
<tr>
<td>Denaturing Histrap Buffer A</td>
<td>20 mM Tris-HCl pH 7.5, 100 mM NaCl, 7 M Urea</td>
<td>His Trap Chromatography for Histones</td>
</tr>
<tr>
<td>Denaturing Histrap Buffer B</td>
<td>20 mM Tris-HCl pH 7.5, 100 mM NaCl, 7 M Urea, 500 mM Imidazole</td>
<td>His Trap Chromatography for Histones</td>
</tr>
<tr>
<td>Denaturing Ion Exchange Buffer A</td>
<td>20 mM Tris-HCl pH 7.5, 0.1 EDTA, DTT, 7 M Urea</td>
<td>Ion exchange Chromatography for Histones</td>
</tr>
<tr>
<td>Denaturing Ion Exchange Buffer B</td>
<td>20 mM Tris-HCl pH 7.5, 0.1 EDTA, DTT, 7 M Urea, 2 M NaCl</td>
<td>Ion exchange Chromatography for Histones</td>
</tr>
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<td>DNA Purification Buffer A</td>
<td>20 mM Tris-HCl pH 7.5, 0.1 EDTA</td>
<td>Protein purification of DNA</td>
</tr>
<tr>
<td>DNA Purification Buffer B</td>
<td>20 mM Tris-HCl pH 7.5, 0.1 EDTA, 2 M NaCl</td>
<td>Protein purification of DNA</td>
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<td>Histone Octamer Refolding Buffer</td>
<td>20 mM Tris-HCl pH 7.5, 0.1 EDTA, 2 M NaCl</td>
<td>Histone Octamer Preparation for Gel filtration and CGC/FPLC</td>
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<tr>
<td>Histone Unfolding Buffer</td>
<td>20 mM Tris-HCl pH 7.5, 10 mM DTT, 7 M Guanidine HCl</td>
<td>Histone Octamer Preparation for Gel filtration and CGC/FPLC</td>
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<tr>
<td>Histone Wash Buffer</td>
<td>50 mM Tris-HCl pH 7.5, 10 mM DTT, 100 mM NaCl, 1 mM EDTA, 1 mM Benzamidine</td>
<td>Protein Extraction and Inclusion Bodies Preparation</td>
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<tr>
<td>Histone Wash Buffer with Triton X-100 1%</td>
<td>51 mM Tris-HCl pH 7.5, 10 mM DTT, 100 mM NaCl, 1 mM EDTA, 1 mM Benzamidine, 1% v/v Triton X-100</td>
<td>Inclusion Body Preparation</td>
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<tr>
<td><strong>Kanamycin</strong></td>
<td><strong>Stock Solution:</strong> 30 mg/ml in ddH₂O. Used at 30 µg/ml</td>
<td><strong>Selection Antibiotic Resistance</strong></td>
</tr>
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<td>----------------------------------------------------------</td>
<td>-------------------------------------</td>
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<tr>
<td><strong>Nucleosome Dialysis Buffer A</strong></td>
<td>0.8 M NaCl, 10mM Tris-HCl pH 7.5, 1mM DTT</td>
<td>Nucleosome Stepwise Dialysis</td>
</tr>
<tr>
<td><strong>Nucleosome Dialysis Buffer B</strong></td>
<td>0.6 M NaCl, 10mM Tris-HCl pH 7.5, 1mM DTT</td>
<td>Nucleosome Stepwise Dialysis</td>
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<tr>
<td><strong>Nucleosome Dialysis Buffer C</strong></td>
<td>0.5 M NaCl, 10mM Tris-HCl pH 7.5, 1mM DTT</td>
<td>Nucleosome Stepwise Dialysis</td>
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<tr>
<td><strong>Nucleosome Dialysis Buffer D</strong></td>
<td>0.1 M NaCl, 10mM Tris-HCl pH 7.5, 1mM DTT</td>
<td>Nucleosome Stepwise Dialysis</td>
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<td><strong>SDS Loading Buffer 6x</strong></td>
<td>125 mM Tris-HCl pH 6.8, 2% v/v of 20% SDS, 60% v/v of 100% Glycerol, DTT, 0.8 mg/ml Bromophenol Blue</td>
<td>SDS-PAGE</td>
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<tr>
<td><strong>Soluble His Trap buffer A</strong></td>
<td>20 mM Tris-HCl pH 7.5, 100 mM NaCl</td>
<td>His Trap Chromatography for Histones</td>
</tr>
<tr>
<td><strong>Soluble His Trap buffer B</strong></td>
<td>20 mM Tris-HCl pH 7.5, 100 mM NaCl, 500 Imidazole</td>
<td>His Trap Chromatography for Histones</td>
</tr>
<tr>
<td><strong>Soluble Ion Exchange Buffer A</strong></td>
<td>20 mM Tris-HCl pH 7.5, 50 mM NaCl</td>
<td>Histone column filtration</td>
</tr>
<tr>
<td><strong>Soluble Ion Exchange Buffer B</strong></td>
<td>20 mM Tris-HCl pH 7.5, 1 M NaCl</td>
<td>Histone column filtration</td>
</tr>
<tr>
<td><strong>Sucrose Loading Buffer 6x</strong></td>
<td>30% v/v sucrose in deionized H₂O</td>
<td>Loading Sample in Native PAGE</td>
</tr>
<tr>
<td><strong>Terrific Broth (TB)</strong></td>
<td>12 g/l Tryptone, 24 g/l Yeast extract, 9.4 g/l KH₂PO₄, 4ml/l Glycerol</td>
<td>Media for Histone Expression</td>
</tr>
<tr>
<td><strong>Wash Buffer</strong></td>
<td>50 mM Tris-HCl pH 7.5, 100 mM NaCl</td>
<td>Preparation for Inclusion Bodies</td>
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<tr>
<td><strong>2YT Broth</strong></td>
<td>16 g/l Tryptone, 10 g/l Yeast Extract, 5 g/l NaCl</td>
<td>Media for Histone Expression</td>
</tr>
</tbody>
</table>

*Table 1. Solutions and Buffers*
2.2. Methods

2.2.1. DNA methods

2.2.1.1. Transformation

50 µl of E.coli competent cells (could be Rosetta2, Star pRL, BL21 PlysS) were stodded on ice with 1 µl of DNA on ice for 30 min. Then a heat shock at 42°C for 60 sec. and placed back to ice for another 10 min. 700 µl of LB were added to the tube and then was incubated at 37°C, 180 rpm, 45 min. 150 µl of this culture were plated in agar plates with resistance antibiotics selection (ampicillin, kanamycin, and chloramphenicol) and incubated over night at 37°C.

2.2.1.2. PCR Mutagenesis

The complementary primers with the single base mutation were designed with the flanking region on each side with an extension of 20 bp with a Tm of 50°C. The plasmid was amplified using the primers with the conditions and reagents of the table 2 and 3. The same reaction with no primers was set as negative control. The template and the newly synthetized strands in the PCR product can be distinguished by their mutilation status basing on that, 1,5µL of DpnI was directly added to each PCR reaction and incubated for 16h at 37°C. Once digested both samples were transformed in Rossetta2 competent cells (as described in 2.2.1 before) and plated in different agar plates. Depending on the number of colonies of the PCR with primers and without primers would validate sadistically if the mutation were inserted or no. if the number on the PCR with primers is significantly bigger than the other, a single colony of that plate will be chose to do a miniprep (Thermo Scientific) and sending it for sequencing to double check if the single mutation is correctly inserted in the plasmid.

<table>
<thead>
<tr>
<th>Description</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA</td>
<td>10 ng</td>
</tr>
<tr>
<td>10 KOD buffer</td>
<td>1 x</td>
</tr>
<tr>
<td>2mM dNTPs</td>
<td>0.2 mM</td>
</tr>
<tr>
<td>25mM Mg2+</td>
<td>2 mM</td>
</tr>
<tr>
<td>10 µM Primer 1</td>
<td>0.3 µM</td>
</tr>
<tr>
<td>10 µM Primer 2</td>
<td>0.3 µM</td>
</tr>
<tr>
<td>KOD polymerase</td>
<td>1U</td>
</tr>
<tr>
<td>water</td>
<td>to adjust volume</td>
</tr>
<tr>
<td>Total volume</td>
<td>50 µL</td>
</tr>
</tbody>
</table>

*Table 2.* PCR conditions for KOD hot start polymerase.
<table>
<thead>
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<th>Step</th>
<th>Temperature (ºC)</th>
<th>Time (min:sec)</th>
<th>Cycles</th>
</tr>
</thead>
<tbody>
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<td>1</td>
</tr>
<tr>
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<td>0:30</td>
<td></td>
</tr>
<tr>
<td>Annealing</td>
<td>56</td>
<td>0:20</td>
<td>16</td>
</tr>
<tr>
<td>Extension</td>
<td>68</td>
<td>0:30/kb</td>
<td></td>
</tr>
<tr>
<td>Final Extension</td>
<td>72</td>
<td>5:00</td>
<td>1</td>
</tr>
<tr>
<td>Hold</td>
<td>4</td>
<td>-</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. PCR program using KOD hot start polymerase.

2.2.1.3. Plasmids

Histone genes in were in the pDS4-SR or in the pET3 plasmid vectors, as shown in the table 4.

<table>
<thead>
<tr>
<th>Vector</th>
<th>Gene insert</th>
<th>Gap between RBS and Gene sequence</th>
<th>De novo synthesis</th>
<th>Antibiotic Resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>pD451-SR</td>
<td>Ct H2AX</td>
<td>No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pD451-SR</td>
<td>Ct H2B</td>
<td>No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pD451-SR</td>
<td>Ct H3</td>
<td>No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pD451-SR</td>
<td>Ct H4</td>
<td>No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pD451-SR</td>
<td>Nc H2A</td>
<td>No</td>
<td>Yes</td>
<td>Kanamycin</td>
</tr>
<tr>
<td>pD451-SR</td>
<td>Nc H2B</td>
<td>No</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pEt3a</td>
<td>Nc H3</td>
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<td></td>
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<td>pEt3a</td>
<td>Hs/Nc H4</td>
<td>Yes</td>
<td></td>
<td>Ampicillin</td>
</tr>
</tbody>
</table>

Table 4. Description of the vector with the correspondent histone gene

2.2.2. Protein methods

2.2.2.1. Histone test expression

Transformed competent cells with the plasmids containing the histones sequences were plated on antibiotic selection plates of agar overnight. Next day a swipe of the colonies was inoculated into 20 ml of fresh media at 37ºC, 180 rpm, approx. 3-4h until the OD was between 0.6-0.8. Then IPTG was added to a final concentration of 0.4 mM. Samples can be taken at 30 min, 2h, 4h, and overnight since that point to run a SDS-PAGE with them.

2.2.2.2. Histone large expression

A swipe of colonies were inoculated in a 2L flask containing 1L of fresh media. The culture was incubated at 37ºC, 180 rpm, approx. 3-4h when the OD was between 0.6-0.8, 1 ml of the sample was taken to run a SDS-PAGE, IPTG was added to a final concentration of 0.4 mM, let the culture incubate for 2-3h more and collect another 1ml of the sample for the SDS-PAGE gel to check if the protein induction expression worked. The media was collected and centrifuged to obtain the pellet that was stored at 20 ºC until further use. The samples collected to check
the protein expression, were aswell spun down to a pellet and resuspended in protein gel loading buffer (PGLB) and heated at 95°C for 5 minutes. SDS-PAGE was run with the samples and standard in Coomassie blue stain for 2 hours. And destained in Coomassie Destain after.

**Equipment used in histone expression:**

- Centrifuge Avanti J-26 XP Beckman Coulter
- Rotor JA17 and JLA 10.5 Beckman Coulter
- Shaker incubator Innova 44 New Brunswick Scientific
- Sonicador. ULTRATURRAX Branson sonifier

2.2.2.3. Histone purification

2.2.2.3.1. Inclusion body preparation

The first step for histone purification is inclusion body preparation. Cell pellet containing histone inclusion bodies were resuspended in Wash Buffer and sonicated on ice bath for 2 minutes at 40% amplitude using Branson sonifier. Cell suspension was then transferred to a 50mL NALGENE tube and centrifuged for 15 minutes at 4°C with 23 000 g. The pellet was retained and washed again with Wash Buffer and centrifuged.

Wash steps were repeated twice with Triton Wash Buffer and finally twice again with Wash Buffer. Inclusion bodies were then solubilized by incubating in 0.5mL DMSO at room temperature on a roller for half an hour followed by 1 h incubation in 5mL Unfolding Buffer.

When pellet was almost dissolved the suspension was centrifuged for 20 minutes with 35 000 G. The supernatant containing protein was collected in a new 50mL tube and diluted to 1 in 20 with histone binding buffer to purify with ion exchange chromatography.

2.2.2.3.2. Cation exchange chromatography

The histone solution, after inclusion body preparation, was centrifuged to remove E. coli DNA contamination or any undissolved matter and filtered through 0.2μm PVDF filter. Protein solution was loaded on Fast flow liquid chromatograph (FPLC) with histone binding buffer using HiTrap SP FF 5mL column with flow 1mL/min-1 and eluted in Elution Buffer.

2.2.2.3.3. Dialysis and Lyophilization

Histones were dialyzed against water to eliminate all the urea containing in the solution. 3h 3 times and then over nigh. Dialysis was performed at 4°C against four changes of refolding buffer. For 1 mg of each histone, dialysis was carried out first in 350mL of refolding buffer twice for 1h each followed by 400mL for 2 h. The result of the dialysis was frozen at -80°C over night and then lyophilized for one or two day until was dry.
2.2.2.4. Concentration determination

Concentration of purified protein was measured with UV spectrophotometer. Purified protein solution was either transferred to a cuvette or measured on a nanodrop pedestal. Absorbance at 276nm (A\textsubscript{276}) measured and purity of the protein was determined by 260:280 ratio in a Thermoscientific spectrophotometer. The absorbance values were noted to calculate accurate concentration using the formula:

\[
C (\text{mgmL}^{-1}) = \frac{[A_{276}]}{\varepsilon} \times \text{MW}
\]

\[
C (\mu\text{M}) = \frac{[A_{276}]}{\varepsilon} \times 10^6
\]

*Concentration (C) determination of pure protein. Where molecular weight (MW) is in Dalton and molar extinction coefficient (\varepsilon) is in M\textsuperscript{-1}/cm\textsuperscript{-1}.*

2.2.2.5. Histone complex refolding

Histones were refolded into dimer, tetramer or octamers using salt-dialysis method. The lyophilized histone aliquots were dissolved in unfolding buffer to a final concentration of 2 mg/mL\textsuperscript{-1}. Unfolding was carried out in unfolding buffer at room temperature for 30 minutes. Concentration of each histone was determined by UV spectrophotometer. The four histone proteins were mixed in an equimolar ratios and volume was adjusted to a final protein concentration of 1 mg/mL\textsuperscript{-1} using unfolding buffer. Histone mixture was then transferred into dialysis bags with 8,000 MW cut off. Dialysis was performed at 4ºC against four changes of refolding buffer. For 1 mg of each histone, dialysis was carried out first in 350 mL of refolding buffer twice for 1 h each followed by 400 mL for 2 h. The final dialysis step was carried out overnight in 1 L refolding buffer. Precipitated protein was removed by centrifugation and concentrated to a final volume of 1 mL. This concentrated sample was then purified by gel filtration chromatography. Refolded histone complexes were always kept at 4ºC. Histone proteins were refolded into octamer using refolding buffer containing 2 M NaCl and histones dimer or tetramer were assembled in refolding buffer with 1 M NaCl.

2.2.2.6. Gel filtration chromatography

Purification of histone complexes were carried out on a Superdex-200 gel filtration column with 0.5 mL flow rate. Column was pre-equilibrated with filtered and degassed refolding buffer. Concentrated histone octamer sample of volume 0.5 mL were loaded into the FPLC system using 1 mL syringe. Purified octamer, tetramer and dimer peaks were collected in 96 well plates with a set fraction size of 0.5 mL. High molecular weight aggregates elutes first at about 7 mL followed by histone octamer at 12 mL to 13 mL and histone dimer at 15 mL for N.
Coriiceps. The purity and stoichiometry of fractions were checked on 15% SDS PAGE. Fractions of equimolar quantity of the histone proteins were pooled and stored at 4ºC.

2.2.3. Nucleosome methods

2.2.3.1. Salt dialysis

Purified octamer was mixed with equal molar amounts of DNA cy-labelled in a solution containing final concentration of 2 M NaCl and 10 mM Tris-Cl, pH 7.5 buffer. The reaction mix was transferred into mini dialysis blocks (Figure) after 30 minutes incubation on ice followed by exchange of buffers for 2 h each in Nucleosome Dialysis Buffer A 0.8 M NaCl, Nucleosome Dialysis Buffer B 0.6 M NaCl, Nucleosome Dialysis Buffer C 0.5 M NaCl and Nucleosome Dialysis Buffer D 0.1 M NaCl at 4ºC (Figure 2). Nucleosomes were stored at 4ºC.

![Diagram of dialysis process](image)

**Figure 5.** Nucleosome dialysis (Doran, 2013). The DNA and histone octamer are placed into the sample chamber in 25 µl volumes and this one is then placed into the pre-chilled dialysis chamber which contains 8 ml of 4ºC dialysis buffer. The assembled unit is then placed on a magnetic stirrer and left to dialyze in a 4ºC cold room.

2.2.3.2. Native gel electrophoresis

A 6% native polyacrylamide gel was prepared and was left to set for 1 h before pre-equilibrating for 3 h at 300 V in 0.2xTBE. From a 100 pmol nucleosome reaction, 2 µL was mixed with 8 µL of 5% sucrose in a 1.5 mL tube on ice. This mixture was loaded on the pre-equilibrated native gel and was further run for 3 h at 300 V. The gel was scanned using fluorescent imager for Cy-labelled samples.
3. RESULTS

3.1. Establishing novel histone expression

The first objective that is required for study the nucleosome of the divergent thermal organisms is to express a significant amount of protein for each histone. To achieve this purpose, proteins were expressed in *E. coli*, where they are expected to be free of post-translational modifications with high expression levels in inclusion bodies that should limit the proteolytic degradation.

3.1.1. De novo sequence optimization

The genes encoding each histone were designed. As no genomic material from the divergent organisms was available, the sequences were synthesized de novo using algorithms optimized for maximal production in *E. coli*. Each gene was optimized for codon usage and to minimize 5' mRNA secondary structure. It is important to achieve this optimization of gene sequence to suit the codon usage of the heterologous host because certain codons that are rare in *E. coli* affect recombinant protein expression (Kane, 1995). Codon optimization also plays an important role to avoid tRNA depletion by eliminating repetitive codons in the genes (Henaut and Danchin, 1996).

3.1.2. Plasmid construction

The chosen vector was pD451-SR (Figure 6) from the company DNA 2.0. This plasmid includes a gene coding for kanamycin resistance (NPT II/Neo) to enable selection. Secondly it contains, the lacI regulatory region of the lac operon that codes for the repressor that binds very tightly to a short DNA sequence just downstream of the promoter near the beginning of lacZ called the lac operator. The repressor binding to the operator interferes with binding of RNAP to the promoter, and therefore transcription occurs only at very low levels. Thirdly, the LacO regulatory region of the lac operon is included. If lactose is missing from the growth medium, the repressor binds very tightly to a short DNA sequence just downstream of the promoter near the beginning of lacZ called the lac operator. The repressor binding to the operator interferes with binding of RNAP to the promoter, and therefore transcription occurs only at very low levels. When cells are grown in the presence of lactose, a lactose metabolite called allolactose, which is a combination of glucose and galactose, binds to the repressor, causing a change in its shape. Thus altered, the repressor is unable to bind to the operator, allowing RNAP to transcribe and thereby leading to higher levels of the encoded proteins. Silencing of the promoter prior to IPTG induction is achieved using symmetrical lac operators (Sadler et al., 1983) spaced around the promoter to maximize cooperativity (Oehler et al., 1994). This operator pair ensures significantly tighter repression than regular lac operators (Lanzer and...
Bujard, 1988). Fourthly, the plasmid contains a mutated form of origin derived from *E. coli* plasmid pBR322 which allows production of greater than 500 copies of plasmid per cell. Fifthly, the promoter T7 ahead of the gene interest is derived strong promoter which is recognized by T7 RNA polymerase. The promoter is controlled by a lac operator sequence that allows induction by addition of IPTG. Finally, a strong ribosome binding site (RBS) is included as a sequence on mRNA that is bound by the ribosome during protein translation. Prokaryotic ribosomes recognize RBSs primarily via base-pairing between the RBS and an unstructured end of the 16s rRNA molecule that forms part of the ribosome. Translation initiation rate of a particular mRNA can be regulated by sequence of the RBS, leading to varying strength (strong, medium or weak).

The histones genes which were introduced into this plasmid were *C. thermophilum* H2AX, H2B, H3, H4, and *N. coriiceps* H2A, H2B. *N. coriiceps* H4 is identical to *H. sapiens* H4.

*Figure 6. Structure of the plasmid with the histone gene CtH2AX.*
3.1.3. *N. coriiceps* H3 mutagenesis

*N. coriiceps* H3 is highly homologous to human histones so this gene was not synthesized de novo, but instead the H3 gene was generated by mutagenesis from human sequences already available in the plasmid pET3a. Mutagenesis primers for the two amino acid changes were designed from the *H. sapiens* H3 sequence to generate the *N. coriiceps* H3 encoding gene (Figure 2). The mutagenesis was successful as the plate with the negative control had 2 colonies and the positive one had 15. The mutagenesis was checked by sequencing the plasmid.

**Figure 7.** *N. coriiceps* H3 mutagenesis sequences. (A) Alignment of protein sequences *H. sapiens* H3 against N. coriiceps cH3. (B) Alignment DNA sequence of HsH3 against NcH3. In black is the two base mutation, in red primer design.
3.2. Optimization of protein expression

All eight plasmids encoding the four histones from the two organisms were transformed into Rosetta2(DE3) pLysS cells. This *E. coli* strain express T7 lysozyme that inhibits the basal activity of T7 RNA polymerase and contributes for lower gene toxicity (Studier, 1991). Growth conditions were at 37°C with 180 rpm agitation. The *E. coli* cells were induced to express the proteins when the cell density reached OD₆₀₀ 0.6-0.8.

3.2.1. Expression media trial

Three different growth media were tested with Rosetta2 to see if a change in media components could enhance the expression. Histones were expressed using 2YT, Terrific Broth and Miller media (Figure 8).

*Figure 8. Expression media optimization. SDS-Gel showing test expression using E. coli strains Rossetta2 in different media: (A) Miller (B) Terrific broth (C) 2YT media showing induction with IPTG (OD 0.6-0.8). Lanes showing uninduced (-) and induced (+). However, expression levels for human H4 (HsH4) were comparatively high.*

Most of the histones of the two divergent thermal organisms demonstrated almost the same moderate expression in *E. coli* after our media optimization strategy, whereas *C. thermophilum* H3 and H4 expressed poorly in all cases. This suggests that media choice does not make a large difference to histone production in any of our cases.

3.2.2. Expression strain trial

Different *E. coli* strains were tested in order to improve the histone expression levels. Although our sequences were already codon optimized, Rossetta 2(DE3) pLysS contains genetically incorporated tRNAs genes for 7 rare codons whereas the Star pRIL strain codes for only 3. The Star pRIL strain also contains a mutation in the gene encoding RNaseE to enhance mRNA stability (Lopez et al., 1999). The pLysS plasmid carries the gene encoding T7 lysozyme and does not provide codon supplementation.
We compared expressions of each histone in Rosetta2 pLysS, BL21(DE3) pLysS and Star pRIL competent cells using 2YT media. The expression levels in Star pRIL and BL21(DE3) didn’t improve significantly for any histone compared to Rosetta2 (Figure 9).

![Image](image.png)

**Figure 9.** Test expression and media optimization. SDS-Gel showing test expression using E. coli strains (A) BL21 pLysS (B) Star pRIL (C) Rossetta2 in 2YT media showing 4 h induction. Lanes showing uninduced (-) and induced (+) cultures with IPTG for H2A, H2B, H3 and H4.

<table>
<thead>
<tr>
<th>Organism</th>
<th>Histone</th>
<th>Gene sequence precedence</th>
<th>Protein lengths</th>
<th>Molecular weight (MW)</th>
<th>Extinction coefficient ($\varepsilon$)</th>
<th>Test Expression</th>
<th>Production Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>N. coriceps</td>
<td>H2A</td>
<td>De novo Synthesized</td>
<td>127</td>
<td>13663,98</td>
<td>4470</td>
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<td>+</td>
</tr>
<tr>
<td></td>
<td>H2B</td>
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<td>125</td>
<td>13760,07</td>
<td>7450</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>H3</td>
<td>Mut. from HsH3</td>
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<td>15404,02</td>
<td>4470</td>
<td>+</td>
<td>++</td>
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<tr>
<td></td>
<td>H4</td>
<td>Same as Hs H4</td>
<td>103</td>
<td>11381,37</td>
<td>5960</td>
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<td>++</td>
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<td>15089,31</td>
<td>7450</td>
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<td>103</td>
<td>11386,34</td>
<td>5960</td>
<td>+</td>
<td>-</td>
</tr>
</tbody>
</table>

**Table 5.** Summary table of origin, state, properties and production of each histone.
3.2.3. Variation in expression between inoculums

Expression levels can sometimes vary from one colony to another. An experiment was designed to compare inoculums derived from 13 separate single colonies of *C. thermophilum* H3 grown independently for expression. All 13 single colony showed equivalent expression (Figure 10).

![Expression trial](image)

*Figure 10. Expression trial. 13 single colonies picked from the same plate of *C. thermophilum* H3. In the 14th line is the + control with the *N. coriiceps* H3.*

3.2.4. Optimization of *C. thermophilum* H4 ribosome binding site

We hypothesized that the reason for some cases of poor expression could be in the structure of the plasmid because noticed that there was no space between the RBS and the sequence of the protein (Figure 11). Other histone expression plasmids with much higher expression for proteins of the same weight and complexity have spaces of at least 6 bases from the RBS to the gene sequence. By reducing mRNA folding free energy near the RBS the rates of translation initiation can dominate expression levels of the histone genes (Kudla et al., 2009; de Smit and Van Duin, 1990). Modifying the stability and folding of mRNA secondary structure near the ribosomal binding site by introducing a sequence with reduced GC content favors translation and quantitatively affects expression levels of recombinant proteins (Kudla et al., 2009; Wu et al., 2004).

Mutagenesis PCR was designed for the plasmid of *C. thermophilum* H4 for insert a 7 base pairs (ATATACAT) spacer between the RBS and the ATG codon.
**Forward:**

AAATAATTTTGTTTAACTTTTTGAGACCTT AAGGAGTAAAAAATATACATATGACTGGTCGTTGTAAGG

**Reverse:**

CCTTACCAGCAGTCATATGTATATTTTTACCTCTTTAAGGTCTAAAAAAGTTAAACAAAAATTATT

---

**Figure 11.** Plasmid structure marking the region where the insertion of the gap and primers design for *C. thermophillum* H4 mutagenesis. Flanking sequences underlined about 20 bp in each side (Tm: 50°C), in red the gap insertion, in blue the RBS.

The mutagenesis was checked by sequencing. We then tested the expression of the new plasmid after the mutagenesis (Figure 12), and observed an increment of the *C. thermophillum* H4 production that was sufficient to proceed with further steps.
Figure 12. SDS PAGE gel comparing expression of the C. thermophilum H4. The result of the PCR mutagenesis of C. thermophilum H4 in lane 2 (+). In lane 1, negative control (-) of C. thermophilum H4 before the mutagenesis. We can appreciate in same growth conditions, media and E. coli strains, how the mutagenesis make an improvement in the expression of the protein.

3.3. Preparation of histones

3.3.1. Large scale expression
Recombinant histones proteins were expressed in 2 L flask with 1 L media and selection antibiotics chloramphenicol (34 µg/ml) for pLysS together with kanamycin (30 µg/ml) for the expression plasmid pD451-SR or ampicillin (100 µg/ml) for pET3a. After growth and induction by IPTG this media was centrifuged in order to obtain the cell pellet with protein in inclusion bodies due to hydrophobic regions that promote the aggregation of the expressed histones at high concentrations (Kane and Hartley, 1988).

3.3.2. Purification by chromatography
The inclusion bodies were lysed by sonication, and washed by centrifugation then solubilized using guanidine hydrochloride due to the positive net charge of histones. Histones were then purified by cation exchange chromatography using a NaCl gradient for elution. All histones showed a similar behavior, with the most ideal case overall being N. coriiceps H3 (Figure 13). SDS PAGE showed that all fractions correspond with the same purity as the chromatogram peak (Figure 6).
Figure 13. Anion exchange chromatography for N. coriiceps H3. (A) Chromatogram of the Cation exchange HiTrap SP FF 5mL GE Healthcare column showing in blue the 280Abs referring to the protein concentration in the fractions. (B) Validation SDS PAGE with the fractions of the pure protein indicated in the chromatogram.

Proteins were analyzed by UV spectrophotometry to calculate the 260:280 nm ratio of absorbance, and by SDS-PAGE. The 260:280 nm ratio was used to check for DNA and RNA contamination. An ideal 260:280 ratio value is in the range 0.5 to 0.8 for proteins with little or no DNA contamination. Above this ratio there is excess nucleic acid contamination which leads to subsequent complications because histones are poorly UV absorbing DNA binding proteins. Can interfere in the refolding of the octamers.

3.3.3. Purification by gel filtration chromatography

The expression levels of C. thermophilum H4 after purification were less than 2mg for a 1 L culture. Purification from inclusion bodies was never reproducible to give sufficient quantities for experiments and led to an excessive 260:280 ratio above 1.5 indicating DNA contamination. To remove DNA contamination, an additional purification over a Superdex 200 10/300 gel filtration column was used in a reducing buffer containing 7 M urea (Figure 14). After this step, the purity of C. thermophilum H4 showed an acceptable 260:280 ratio of 0.8.
Figure 14. Process column filtration eliminate DNA. (A) Cation exchange chromatography of C. thermophilum H4, (B) SDS-page gel validating the protein in the fractions with high rate of DNA contamination (C) DNA contamination which was removed by gel filtration chromatography using urea buffer after cation exchange chromatography purification Superdex 200 10/300 GL column. (D) SDS-page gel validating the protein with free-DNA removal filtration column. Quality controls measures included, apart SDS-PAGE gel, the UV spectrophotometry.

After purification, histones were dialyzed at 4ºC to eliminate all urea and salt. The dialysate was frozen at -80ºC over night, proteins were then lyophilized in individuals tubes for long term storage at room temperature.

3.4. Assembly of octamers refolded from purified histones

The octamer refolding property of histones was compared for histones from two thermal organisms by mixing equal molar amounts of H2A, H2B, H3 and H4 after they were completely unfolded in 7M Guanidium. This mix was then dialyzed against 3 changes of 2M NaCl buffer to
fold histones into octamers (Luger et al., 1997b). It is in 2M NaCl where the histone octamer complex is in the most stable form because it is dominated by the hydrophobic interactions between the histones and by potential cationic repulsions of the highly basic proteins (Arents et al., 1991; Luger et al., 1997a). Octamer complexes of histones were separated from aggregates and other partial complexes using a Superdex 200 10/300 column. A typical chromatogram of octamers shows elution high molecular weight aggregates at 7 mL followed by octamers complexes at 12 mL to 13 mL, and H2A-H2B dimer peak at 15 mL (Figure 7A).

3.4.1. *N. coriiceps* octamers

*N. coriiceps* octamer assembly was very successful by mixing in equimolar quantities all four histones and using the gel filtration (Figure 15A) to obtain pure octamer fractions (Figure 15B). The fractions were checked by SDS PAGE (Fig 15C) for the nucleosome assembly. Octamers could be stored at 4°C for several weeks.

![Figure 15](image-url)

**Figure 15.** *N. coriiceps* histones refolded into octamers. (A) Chromatogram with the *N. coriiceps* octamers by column filtration Superdex 200 10/300 GL column. (B) SDS PAGE gel with GFC fractions confirming the presence of individual histones and their stoichiometry corresponding to the octamer peak at 12mL elution volume. The composition of the four histones in this octamer complex is qualitatively proportional as expected resulting in perfect yield of histone octamer. (C) Nucleosomes of *N. coriiceps* and *X. laevis*. 
3.4.2. C. thermophilum histones do not refold into octamers

The peak elution fraction resulting from octamer gel filtration is 11-12 ml, but this zone did not show any octamers by SDS PAGE for C. thermophilum refolding (Figure 8A). This suggests that the small peak at 11 ml corresponds to a complex different than octamer, possibly due to limited aggregation of H3, H4 or H2A, H2B histones or from DNA (Figure 16). A reason for the problem of C. thermophilum not forming octamers could be the high contamination of DNA in the histones production making it very difficult to perform exact quantification by UV absorbance of the amount protein and secondly, difficult the histone interaction and assembly into octamers in all our attempts. We were unable to refold C. thermophilum octamers. Consequently, C. thermophilum nucleosomes could not be formed. The inability of C. thermophilum histones to form octamers could maybe show that the histone octamer assembly must be mediated by DNA sequence as C. thermophilum histones are unable to refold into stable tetramers and octamers in absence of DNA by standard salt dialysis method.

Figure 16. C. thermophilum histones do not refold in octamers. (A) Chromatogram with the N. coriceps octamers by column filtration Superdex 200 10/300 GL column. The GFC analysis shows a very small peak at 11ml that corresponds to a structure larger than the octamer, suggesting an inability of C. thermophilum to form stable octamers. Large aggregates elute first at 7ml followed by dimers and tetramers form 16ml. (B) C. thermophilum histone refolding almost entirely resulted in precipitation; we can appreciate two bands in Lanes 1 and 2 from the raw sample before load to the GFC. The rest of the SDS Page could not validate the other peaks that were registered in the chromatogram. The occurrence of these aggregates and absence of any octamer peak were consistent in all trials despite many refolding attempts.

3.5. Nucleosome assembly

Nucleosomes can be assembled in vitro using histone octamer complexes along with a DNA sequence of 147 bp or greater by salt gradient dialysis. We have used the very well characterized nucleosome positioning DNA sequence 54A54 (255 bp). Purified octamers were mixed with DNA at 2 M NaCl and then sequentially dialyzed in reducing salt concentrations to physiologically relevant level 0.1 M NaCl. When the salt concentration is reduced to 1 M, the
octamer dissociates into (H3–H4)2 tetramer and two H2A–H2B dimers, and then at 0.85 M NaCl DNA binds to the tetramer, and at 0.65 M NaCl, H2A–H2B dimers bind on either side of the tetramer to form the nucleosome (Yager, 1989; Germond et al., 1976). The reason to choose this method is that it is highly efficient to avoid non-specific histone–DNA aggregation when mixing pure octamer with DNA.

3.5.1. *N. coriiceps* nucleosomes in vitro by salt dialysis

The ability of *N. coriiceps* octamers to form nucleosomes on 147 bp of the 54A54 sequence of DNA was tested. Purified octamer fractions from gel filtration chromatography of *N. coriiceps* and control *X. laevis* were mixed with DNA at 1:1 ratio and dialyzed by reducing the salt concentration. Nucleosome samples were then validated by native PAGE (Figure 17). Native gel interpretations of nucleosome positioning are not always obvious because the flexibility of DNA at entry/exit angles is variable (Pennings et al., 1991).

![N. coriiceps Nucleosomes and tetrasomes on 54A54 DNA sequence. Native PAGE showing migration of tetrasomes and nucleosomes of N. coriiceps and X. laevis on 54A54 fluorescence labeled by salt dialysis. First and second band are for nucleosomes and di-tetrasomes respectively and third band is free-DNA. X. laevis was loaded in Lane 1, N. coriiceps was loaded in Lane 7 and Free-DNA was loaded in Lane 3 as negative control.](image)

The native PAGE of histone assembly with free DNA showed the capability to form nucleosomes with in both cases, although with an excess of unbound DNA. The mobility of the nucleosome bands of *N. coriiceps* equivalent to *X. laevis.*
4. DISCUSSION

4.1. Histone selection and gene design

There is a general assumption that at the chromatin level all eukaryotes package and process their genomes in an equivalent fashion. We wanted to include two divergent thermal organisms that would test eukaryotic adaptation of chromatin to temperature to allow us to investigate the genome packaging mechanism. To compare relevant nucleosome properties, it was important that the genomes of divergent species be completely sequenced to ensure representative canonical histones were used, avoiding divergent isoforms or variants.

The classification of histones is mainly dependent on their amino acid sequence identity with mammals. Alignment of histone sequences from divergent thermal eukaryotes with *H. sapiens* shows that *C. thermophilum* histones are somewhat divergent, with only 65% identity in H2B. There is very high similarity of H3 and H4 between *N. coriiceps* and *H. sapiens*. H4 is widely considered to be the most conserved histone and only differs by a few amino acids in animals, plants and most fungi.

However, there are many differences in the amino acid positions of *C. thermophilum* in H3 and H4 histones that can be potentially critical for formation of histone tetramer interfaces and hence for octamer stability. Histone sequence alignments of divergent eukaryotes to higher eukaryotes showed significant differences in the regions that may be critical for histone-histone interaction in octamer and histone-DNA interactions in nucleosomes.

4.2. Protein expression and purification

The expression of histones is sensitive sequence and to growth conditions in *E. coli*. We were able to create histone genes for our two divergent organisms by using gene optimization calculations and gene synthesis technology. Sufficient amounts of histones were produced for the biochemical assays by recombinant protein expression in *E. coli*, to produce typically 4 mg from a 1,5 L culture. Testing expression in different strains of *E. coli* and using different media did not produce any obvious changes in yield. However, we found that expression of *C. thermophilum* H4 was improved by optimizing the plasmid to suit *E. coli* to ensure that the RBS sequence is correctly placed. Contamination of DNA was always a problem and we always took care, to avoid excess DNA in the protein sample. Purification using gel filtration was necessary to remove DNA in one case.
4.3. Octamers refolding and nucleosome

*C. thermophilum* histones did not form octamers and mostly precipitated, although some histone dimers remained in solution. Inspection of the amino acid sequence of the histones reveals four changes in amino acid sequence in the H4, at A49, S61, S65 and S84 that differ from *N. coriiceps*, *H. sapiens* and *X. laevis*. The mutation of these residues could affect in the interaction of H4 with H3, making the conformation less stable and this disfavoring octamer stability. This could be the reason *C. thermophilum* does not form octamer despite the conservation of the rest of the amino acid sequence.

Native PAGE of *N. coriiceps* histones assembled on 54A54 DNA shows they have the capability to form mono-nucleosomes. There were differences in the mobility of these nucleosome, with *N. coriiceps* bands being more diffuse suggesting they are more dynamic. The diffuse nucleosome bands could also suggest that the DNA is not tightly wrapped around octamer, with DNA less tightly bound at the entry/exit site of the nucleosome affecting hydrodynamic behavior (Flaus et al., 1996).

The inability of *C. thermophilum* histones to form octamers prevented formation of nucleosomes by the normal salt dialysis technique. Instead, a salt-urea dialysis approach to assemble nucleosomes from individual histones could be used in future experiments.
5. CONCLUSIONS

Histone sequences, even though are highly conserved, have significant level of diversity across the Eukaryotic kingdom. This shows that histone evolution may be more complicated than previously assumed and the biochemical properties of divergent histones may be explicable based on structural interactions. We used an efficient method to produce recombinant histones in *E. coli* in large quantities. Heterologous protein expression of histones can be improved by optimizing gene sequences and growth conditions. For most of the histones, genes synthesis with codon adaptation to *E. coli* and altering the 5’ end of mRNA structure lead sufficiently to high level expression. For poorly expressing histones, optimization of growth conditions by testing *E. coli* strains or growth media didn’t show any positive effect on expression levels. It is important for the correct expression of some proteins that a space exists between RBS and the sequence of that protein. A common scale up and purification strategy that could be applied to all the histones to achieve sufficient amounts of protein for biochemical assays. The results from octamer refolding at high ionic strength show differences in complex formation for the two divergent thermal organisms. *N. coriiceps* histones show similar octamer refolding behavior to *X. laevis*. *C. thermophilum* histone largely precipitated and did not form octamers by salt dialysis. Results from nucleosome assembly show that histone octamers from *N. coriiceps* and *X. laevis* were able to assemble into nucleosomes by standard methods of salt dialysis. There was difference in the mobility nucleosomes from *N. coriiceps* and *X. laevis*, demonstrating distinctive hydrodynamic properties of nucleosomes. This suggests some variation in binding of DNA around the octamer either in a static or dynamic mode.
6. APPENDIX

6.1. Histone Genes sequences

_C. thermophilum_

**CtH2AX**:

```
ATGACCGGTGGCAAAAGCGGTGGCAAAGCATCGGGTACAAAATCAGCGCAGTCTCGTTCCTCCAAAGCTGGCCTG
GCCTTCCGCGCCGCGTTCTGGAAATCTCGGCCTCGGAAATCCTCGGAACTCTCGGCTGACAAT
AAGAAAACCCCGTATCTCCGAGCCTTCGAGCAGCTATCCTTGCCCAAGAAGCTGAGCTTAAAAATACGCTGCGC
CATGTGACCATCGCACAGGGCGGTGTACTGCCGAATATCCATCAGAACCTGCTGCCGAAAAAGACTGCTAAGACG
GGAAAAACTTAAACCGCCTGGAAATATACCTGCTGGTAATGCGGCCCGTGACAAT
```

**CtH2B**:

```
ATGGCACCGAAAGCGGCAATGCGCGCAGAAAAAACCGGCATCCAAGGCACCAGCATCTACCGCTAGCAAGGCTCCGTCT
GAAAAAAAGATCGCCGGTAAAAAACTCCGGTACGCGGTTGAAGAAAAAAAAAGTGAACCAAGAGCGTGAAGAAAGA
CCTACAGCTCGTATATTTATAAAGTTCTTAAGCAAGTACATCCGGACACTGGCATCTCTAATCGTGCTATGTCTATC
CTGAATTCATTCCTGTAATGTATATTTCGACGCGTGGCACGGCGGCTAAATCTGACTGCGGAAATCCTGGAGCTGGCTGG
TAATGCTGCTCCGGAAGATTCAACCTTTGTCGGCAATCCCGCTAGCCAAAAGCTGAGACTTACCCAGTACCGTGAA
CTGCAGCCGCTGCTGGAAATATACCTGCTGGTAATGCGGCCCGTGACAAT
```

**CtH3**:

```
ATGGCTCGTACTAAACAGACTGCGCGTAAATCCACCGGTGGCAAAGCTCCTCGCAAACAGCTGGCGTCTAAAGCACGCC
CGACAAATCTGCACCGAGCACTGGTGGCGTGAAAAAACCACACCGCTATAAACCGGGCACGGTGTTAAACGTATTTC
AGCGCTGCTGGAAATATACCTGCTGGTAATGCGGCCCGTGACAAT
```

**CtH4**:

```
ATGACTGTCGTGTTAAAGGTGTTAAGGCTTCTGGTAAAGGGAGGCACCCGCTTGGCAAAAGCTCCGGTGGTTAAAGGTGI
ATCCCCGGCCTGCTGGCAAAAGCGGTGGTAAAGCGCGTGCAAAAGCAAAAACCCGTTCGAGCCGTGCAGGTTTGCA
ATTCCCGGTGGGTCGTGTGCACCGCCATCTGCGCAAAGGCAACTACGCTCACCGAGAGCTGCTGCTGCTGCTGCTGC
ATCTATGGAAGGCACCGCACCGCTGCTGGTAAAGGTGGATGTTGCTATTCGTTAATGCGCCGCTACCCAGAAGAAGGTT
GAATTCATTCCTGGAATATACCTGCTGGTAATGCGGCCCGTGACAAT
```

_N. coriiceps_

**NcH2A**:

```
ATGTCTGGTCGTGGCAAAACCGGTGGTAAATCCACCGGTGGGCAAAAAGCGGTGGTAAAGCGCGTGCAAAAGCAAAAACCC
GTTCGAGCCGTGCAGGTTTGCAATTCCCGGTGGGTCGTGTGCACCGCCATCTGCGCAAAGGCAACTACGCTCACCGAGAGC
TCTGCTGCTGGTAATGCGGCCCGTGACAAT
```

**NcH2B**:

```
ATGCCAGAAGCAGCATCTGCTGGCAAAAGCGGTGGTAAATCCACCGGTGGGCAAAAAGCGGTGGTAAAGCGCGTGCAAAAGC
GTTCGAGCCGTGCAGGTTTGCAATTCCCGGTGGGTCGTGTGCACCGCCATCTGCGCAAAGGCAACTACGCTCACCGAGAGC
TCTGCTGCTGGTAATGCGGCCCGTGACAAT
```

**NcH3**:

```
ATGGCCCGTACCAAGCAGCCGCCGGCTGCAAAAGCGGTGGTAAATCCACCGGTGGGCAAAAAGCGGTGGTAAAGCGCGTG
CACAATCCACCGGTGGGCAAAAAGCGGTGGTAAATCCACCGGTGGGCAAAAAGCGGTGGTAAAGCGCGTGCAAAAGC
GTTCGAGCCGTGCAGGTTTGCAATTCCCGGTGGGTCGTGTGCACCGCCATCTGCGCAAAGGCAACTACGCTCACCGAGAGC
TCTGCTGCTGGTAATGCGGCCCGTGACAAT
```
NcH4/HeH4:
ATGTCTGGTCGTGGTAAAGGTGGCAAAGGTCTGGGTAAAGGTGGTGCGAAACGTCATCGTAAAATCCTGCGCGA
TAACATTTGACGCGCATTACAAACCAACAGCACCTTCCCCGTGGGGTCTGCA
GTTTCGGTGTAGCGTGTCATCTGCCTCTCTGGCAAAAGGCACACTGCAGGAGTGGTTGGTCTGGGTCTGGCTCCTGC
CTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
CGTCACCATTTGCAGGAGGCGGGCTTTTGGCCCAATATCCAGGCGGTCTTCTCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

H. sapiens
HsH2AX:
ATGTCAAGGTCGCGGTAAAACTGGCGGGAAAGCCCGTGCGAAAGCGAAATCGCGCAGTTCCCGTGCCGGTCTGCA
GTTTCCGGTGTAGCGTGTCATCTGCCTCTCTGGCAAAAGGCACACTGCAGGAGTGGTTGGTCTGGGTCTGGCTCCTGC
CTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
AGTCACTATCGTCGAGGGGCGTTCTGCCCCAACACTCCAGGCTTACCTGCTGTTACCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

HsH2A:
ATGTCAAGGTCGCGGTAAAACTGGCGGGAAAGCCCGTGCGAAAGCGAAATCGCGCAGTTCCCGTGCCGGTCTGCA
GTTTCCGGTGTAGCGTGTCATCTGCCTCTCTGGCAAAAGGCACACTGCAGGAGTGGTTGGTCTGGGTCTGGCTCCTGC
CTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
AGTCACTATCGTCGAGGGGCGTTCTGCCCCAACACTCCAGGCTTACCTGCTGTTACCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

HsH3:
ATGGCCCGTACCAAGCAGACCGCCCGGTAAATCCACCGGAGGAAAGGCTCCAAGAACACCGGCCTACCAGGGCATGCATGAC
AGACGGGGAAAGCAAGCGGTGTAAAGGTGGTAAAGGTCTGGGTAAAGGTGGTGCTAAACGTCACCGTAAAGTTCTGGCA
AACATTTGACGCGCATTACAAACCAACAGCACCTTCCCCGTGGGGTCTGCA
GTTTCGGTGTAGCGTGTCATCTGCCTCTCTGGCAAAAGGCACACTGCAGGAGTGGTTGGTCTGGGTCTGGCTCCTGC
CTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
AGTCACTATCGTCGAGGGGCGTTCTGCCCCAACACTCCAGGCTTACCTGCTGTTACCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

X. laevis
XhH2A:
ATGTCAGGGAAGGCGAGAAGGCGATAAAGGCGTGTTAAAGGCGCTAAGGCGCCAAGACTCGGCTTCTCG
GCTGTGGGCTACAGGTTCCTTCTTGCCGCGGCGCGCGTAAAAGCAGGTTACTCGGGCGC
GCTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
AGTCACTATCGTCGAGGGGCGTTCTGCCCCAACACTCCAGGCTTACCTGCTGTTACCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

X. laevis
XhH2B:
ATGCCAAGAACCGGCCAAGTCCGCTCCAGCCCCGAAGAAAGGCTCCAAGAACACCGGCCTACCAGGGCATGCATGAC
AGACGGGGAAAGCAAGCGGTGTAAAGGTGGTAAAGGTCTGGGTAAAGGTGGTGCTAAACGTCACCGTAAAGTTCTGGCA
AACATTTGACGCGCATTACAAACCAACAGCACCTTCCCCGTGGGGTCTGCA
GTTTCGGTGTAGCGTGTCATCTGCCTCTCTGGCAAAAGGCACACTGCAGGAGTGGTTGGTCTGGGTCTGGCTCCTGC
CTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
AGTCACTATCGTCGAGGGGCGTTCTGCCCCAACACTCCAGGCTTACCTGCTGTTACCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

X. laevis
XhH3:
ATGTCAGGGAAGGCGAGAAGGCGATAAAGGCGTGTTAAAGGCGCTAAGGCGCCAAGACTCGGCTTCTCG
GCTGTGGGCTACAGGTTCCTTCTTGCCGCGGCGGCGCGTAAAAGCAGGTTACTCGGGCGC
GCTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
AGTCACTATCGTCGAGGGGCGTTCTGCCCCAACACTCCAGGCTTACCTGCTGTTACCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

X. laevis
XhH4:
ATGTCTGGTCGTGGTAAAGGTGGTAAAGGTCTGGGTAAAGGTGGTGCTAAACGTCACCGTAAAGTTCTGGCA
AACATTTGACGCGCATTACAAACCAACAGCACCTTCCCCGTGGGGTCTGCA
GTTTCGGTGTAGCGTGTCATCTGCCTCTCTGGCAAAAGGCACACTGCAGGAGTGGTTGGTCTGGGTCTGGCTCCTGC
CTACTTAAGCAGGGGTTGGAGTCTGCAGACGGCGAAATCTGGAACGGCTGTAATGCGACGCTGACAAACAA
GAAAACCCGCACTATTCCAGGCGCGATTGCGATTGACGGAAGTCAGCAAAAAACTCTCGGGAG
AGTCACTATCGTCGAGGGGCGTTCTGCCCCAACACTCCAGGCTTACCTGCTGTTACCCCGAAGAAACAGAGGCGTACCTGGA
TGAGCGGAAACGCGCTTTCTGTGCAAGAAGAAGAGCCACAGCGCAGGGTG

X. laevis
CCTCCCGCCTGTATTACAAACAGGGCTTCCGAGGAGGCATTCCGACTGCTGCTGCCTGGGGAGTTGGCCAAACACGCCGTGTCCGAGGGCACCAAGGCTGTCACCAAGTACACCAGCGCCAAG

**XlH3:**
ATGGCCCGTACCAAGCAGACCGCCCGTAAAATCCACCGGAGGGAAGGCTCCCCGCAAGCAGCTGGCCACCAAGGC
AGCCAGGAAGTCCGCTCTGGTCTACCGGCGATCAGAAAGACCTCAGGTTACCGGGCGCCGACAGTCGGCTTCCCG
CGAGATCCGGCCGCTACCAAGAAATCCACCGGAGGTCTGCTACCCGCAAACCTGCTTCCCAGCGCCGTGGTCGAGATGC
GCTCAGAATCTCAACGCGACCTCGGCTCCAGAGCTGGTTATTAGTGTCTGACAGAGGCGAGAGCTTAT
CTGCGTCGCTCTCTTTGAGGACCAACACTGTGCGCCATCCAGCCAGCAGAAGGTCAGTCACCACATCTGCCCGAGAGGAGGCT
AGCTGGCCCGAATCCCGGAGGAGGGCCT

**XlH4:**
ATGTCTGGTCGTGGTAAAGGTGGTAAAGGTCTGGGTAAAGGTGGTGCTAAACGTCAACCGTAAAGATAAGTTCTGGTAC
AACATCCAGGGTATCACCAAGCCGGCTATCCGGTCTGTCGCTGGTGTTGTTAAGCTATCTCCGCTGGA
TCTAGAAGAAACCGGCGGGTGTTCTTTGAGGATACCTGCTTACCGGCCCATCCACGCCG
CGAGATCCGGCCGCTACCAAGAAATCCACCGGAGGTCTGCTACCCGCAAACCTGCTTCCCAGCGCCGTGGTCGAGATGC
GCTCAGAATCTCAACGCGACCTCGGCTCCAGAGCTGGTTATTAGTGTCTGACAGAGGCGAGAGCTTAT
CTGCGTCGCTCTCTTTGAGGACCAACACTGTGCGCCATCCAGCCAGCAGAAGGTCAGTCACCACATCTGCCCGAGAGGAGGCT
AGCTGGCCCGAATCCCGGAGGAGGGCCT

**6.2. Histone Protein sequences**

**C. thermophilum**

**Ch2A:**
MTGGKSGGKAGTSAQ5RS55SKLAFLPGVPRVHRLLRKNGYAQRVGAGPVLYAALVLAAAILELAGNAARDNKT
RIIPRHQLAIRNDEELNKLHGVHTAQTGRLPNHIQNLPLKTTAKTKNLSEL

**Ch2B:**
MAPKADAQKPKPSKAPASTAKPEKSEDAKTKAAEGKKTARKETYSYKYKLVQPVDTPGISNRMASILNSFV
NDIFERVATEASKLAAYNKSTISSREIQTVARLILPELAKHAVSEGTKAVKTYSSSTK

**Ch3:**
MARTKQTARKSTGGKAPPKRLASKAARSAPSTGGVPPKHPYRKPTVALREIRYQKSTELLRTKLFQRLVREIAQDFK
SDLRFQSSAIGALQESVESVLYLFEDNTLCAIHAKRVTIOSEQIDQLARRGRRERN

**Ch4:**
MTGRGKKGGKLGGKAKHRKILRNIQGITKPAIRRLRGGVVKRISAMYPEETRGVLKFSLESFVIRDAVTYEHAKRK
TVTSLDVYALKRQGRTLYFGF

**N. Coriiceps**

**Nh2A:**
MSGRGKTKGGKARAKATKRRASSLQPQFPVPVRHVRLLRNYGNYAHRVGAGAVYLAALVLAAILELAGNAARDNKT
RIIPRHQLAIRNDEELNKLHGVHTAQTGRLPNHIQNLPLKTTAKTKNLSEL

**Nh2B:**
MPEAASVKKPGSKKAVTCKTPSKGKRRKSRKESYAIYVKMVQVHPDTGISSKAMGIMNCVSDIERAGEASRL
AHYNKRSTTSREIQTVARLILPELAKHAVSEGTKAVKTYSSSTK

**Nh3:**
MARTKQTARKSTGGKAPPKRLASKAARSAPSTGGVPPKHPYRKPTVALREIRYQKSTELLRTKLFQRLVREIAQDFK
SDLRFQSSAIGALQESVESVLYLFEDNTLCAIHAKRVTIOSEQIDQLARRGRRERN

**Nh4:**
MSGRGKGGKLGGKAKHRKILRNIQGITKPAIRRLRGGVVKRISAMYPEETRGVLKFSLESFVIRDAVTYEHAKRK
TVTSLDVYALKRQGRTLYFGF

**H. sapiens**

**Hs2A:**
MSGRGKTKGGKARAKATKRRASSLQPQFPVPVRHVRLLRNYGNYAHRVGAGAVYLAALVLAAILELAGNAARDNKT
RIIPRHQLAIRNDEELNKLHGVHTAQTGRLPNHIQNLPLKTTAKTKNLSEL

**Hs2B:**
MSGRGKGGKLGGKAKHRKILRNIQGITKPAIRRLRGGVVKRISAMYPEETRGVLKFSLESFVIRDAVTYEHAKRK
TVTSLDVYALKRQGRTLYFGF
MPEPAKAPAPKGSKKAVTKAQKDGKKRKRKRSKESYVYYKVLKVHQDGISSKAMGIMNSFVNDIFERIAEASRLAHYNKRSTISREIQTAVRLLLPGELAHAVSEGATKVAVKYTEASAK

**HsH3:**
MARTKQTARKSTGGKAPRKQLATKAARKSAPATGGVKKPHRYPGTVLREIRRRYKSTELLIRKLPQRLVREIAQDFKTDLRFQSSAVMALQEACEAYVGLFEDTNCARIEAHAKRVITMPKQQLLRIRRGERA

**HsH4:**
MSGRQKGGKGLGGAKHRKVLVRDNIQQGITKPAIRRLARRGGVKRISGLYETRGLVFLNVIRADVYTHEAKRKTVTAMDVYALKQGRRTLYGFGG

**X. laevis**

**XlH2A:**
MSGRGKQGGKTRAKTRSSRAGLQFPVGRVHRLRKGNYAERVGAGAPVYLAAVLETAEILELAGNAARDNKKTRIIPRHLQAVRNEELNLGVRDIAQQGVLPLQIVLSLPKTEESSKASKK

**XlH2B:**
MAKSAPAPKKGSKAVTKTQKKDGKKRRKTRKRESYAIYVVKLVQHPDTGQSSKAMSIMNSVNDVIFERIAGEASRLAHYNKRSTISREIQTAVRLLLPGELAKHAVSEGATKVAVKYTEASAK

**XlH3:**
MARTKQTARKSTGGKAPRKQLATKAARKSAPATGGVKKPHRYPGTVLREIRRKYKSTELLIRKLPQRLVREIAQDFKTDLRFQSSAVMALQEACEAYVGLFEDTNCARIEAHAKRVITMPKQQLLRIRRGERA

**XlH4:**
MSGRGKGGKGLGGAKHRKVLVRDNIQQGITKPAIRRLARRGGVKRISGLYETRGLVFLNVIRADVYTHEAKRKTVTAMDVYALKQGRRTLYGFGG

6.3. **Histone proteins alignments**
7. REFERENCES


PATWAL, I (2014). Properties of divergent histones. Prof. Andrew Flaus, PhD. Galway: National University of Ireland Galway


