Results and limits of the soliton theory of DNA transcription

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Abstract. It has been suggested by several authors that nonlinear excitations, in particular solitary waves, could play a fundamental functional role in the process of DNA transcription, providing the opening of the double chain needed for RNA Polymerase to be able to copy the genetic code. Some models have been proposed to model the relevant DNA dynamics in terms of a reduced number of effective degrees of freedom. Here I discuss advantages and disadvantages of such an approach, and discuss in more detail one of the models, i.e. the one proposed by Yakushevich.

1. Introduction

The first step in the replication of DNA [33] is its transcription, from the original contained in the cell to a copy – the RNA messenger – which will then be used as a “master copy” for producing actual copies of the genetic information. The evolutionary advantage of such a messenger is obvious: in this way, the original DNA is opened – and thus less protected – for as small a time as possible.

The transcription process is carried out by a specialized enzyme, the RNA Polymerase (RNAP); here we are mostly interested in the dynamical aspect of the transcription, which is roughly speaking as follows. The RNAP opens a “transcription bubble” of a size of about 15-20 base pairs, and then travels along the DNA chain keeping the size of the open region more and less constant, i.e. providing at the same time to open the chain in front of it and closing back the one behind.

The process undergoes several (even quite long in time) stops, but in the active phase the RNAP proceeds along the DNA chain at a speed of 500 – 1000 base pairs per second. Since each base pair is linked by two or three hydrogen bonds, the energy involved in such a process, even considering only the one to open (and close) the DNA chain, is of the order of thousands H bonds per second.

Obviously, as biological systems live at a temperature of the order of 300 Kelvin, thermal energy is widely available, but the problem is how is this energy focused at the right place and with the right timing to operate the process.

Needless to say, the problem of energy transport and focusing is of more general relevance in Biology and biological systems – and one indeed which has to be studied by

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Biological Physics. Thus, the study of this problem in the dynamics of DNA transcription is not only relevant *per se* and for the fundamental role of DNA transcription and then replication in life, but can also have a more general relevance for the study of energy transport in living systems.

It has been suggested by several authors that Nonlinear Dynamics can have a word to say concerning this, and that energy transport could happen by means of nonlinear excitations travelling along biological chains, such as long proteins or indeed DNA.

The most widely known theory in this context is probably that of “Davydov’s soliton” [7,8], which is of essentially quantum nature. Theories concerning nonlinear excitations in DNA chains and their functional role have been proposed by several authors, following the seminal work of Englander, Kallenbach, Heeger, Krumhansl and Litwin [11] (and sometimes it is difficult to understand priorities, so I will not assign any); these present differences, but many more common aspects.

Excellent reviews exist on Nonlinear Waves in biological systems [34,35]; I would also like to mention, as a source of a number of inspiring material on the relations between nonlinear structures and biology (or more specifically biological molecules) the volume collecting the proceedings of the Les Houches 1994 School [28].

My goal here is not to review and/or compare the different theories proposed for these would-be excitations of DNA connected with transcription, but rather discuss the merit and limitations of the general idea of what I will call, with a deliberately ambiguous expression, ”soliton-helped transcription”.

Roughly speaking, the idea is that there are nonlinear excitations travelling along the DNA chains, causing a local opening of the double chain. The RNAP could then travel along with these, and use the opening of the chain to read the DNA sequence without having to focus the energy needed to open the double chain.

Another advantage of this theory is connected with the closing of the double chain after the RNAP has passed: if this is done in a non-coordinated way, it will generate a substantial quantity of random motion and thus of thermal energy; if instead this correspond to the passing of a nonlinear excitation (maybe in this context it is more appropriate to call our soliton a “nonlinear coherent structure”), no thermal energy is generated when the system recovers its local fundamental state.

I will of course also discuss quantitative results and predictions, and not just qualitative ideas; in doing this I will refer to a specific model, the one suggested by L.V. Yakusheevich [39,40] in the late eighties, on which I have done some work and which is well suited to analytical investigation. Needless to say, the very fact that it is amenable to analytic results means that it is an oversimplified model, and it is not difficult to find in the literature – or to build by oneself – more detailed models. However, as I will advocate in more detail in the sequel, I believe the study of such simple models can teach us a lot.

Before proceeding, there is a point that should be made clear for the sake of the more mathematical reader: the nonlinear excitations we wish to consider are not necessarily solitons in the proper mathematical sense [1,9,32], but rather solitary waves; however, it is by now common – and more attractive – to call them solitons, and I will conform to this usage.

Also, the word “soliton” is used in two senses, a topological one [10] together with the
dynamical one [1,9,32]. The solitons to be considered here will be both solitary waves in dynamical sense and $S^1$-topological solitons.

2. What should a model contain?

The DNA molecule is an awfully (or wonderfully) complicated one; each segment corresponding to a base pair has about 100 degrees of freedom, and a DNA molecule can contain $10^{10}$ (or even $10^{13}$ as for salamanders) such segments, thus having $10^{12}$ degrees of freedom.

As if this was not enough, DNA is very sensitive on the “boundary conditions”, i.e. to the characteristics (physical and chemical) of its environment, and no chemist or biochemist would think of DNA “in abstract”, i.e. without specifying in which environment it operates.

With such numbers (even apart from the problem of modelling the interactions of the DNA molecule with its environment) it is clear that there is no hope of detailed mathematical modelling. Actually, these are so large that we are quite justified in applying the thermodynamical limit (i.e. consider an infinite chain), and resorting to statistical mechanics [22,24]; indeed, several approaches have been attempted in this direction, and with some success.

However, the dynamical models which have been proposed to study the dynamics of DNA transcription, only consider an extremely limited number of degrees of freedom (one per base in the Yaushevich model); indeed, it seems hopeless to study a model with more than a very few degrees of freedom per segment and investigate if it has solitons, not to say determine the detailed dynamics of these.

Thus the first question we should ask is: does it make any sense to study such simplified models of such an enormously complicated system as DNA?

Substituting simple mathematical models to complicate real system is, of course, what has always been done in theoretical – and not only theoretical – Physics, and it has been successful many times; but this is of course not a good reason to say that it can be done here as well, not to say that in a biological system the complexity is somehow much more fundamental, inherent to the system, than in Physics, where we can e.g. study gravitation without the disturbances of air by dropping masses in a vacuum tube: in Biology, “simplifying the system” most probably means killing it, and biologists and biochemists are well justified in their diffidence towards “Physics-style” simplified models.

I would like at this point to open a short parenthesis to mention the problem of dissociation (or melting) of DNA: experimental data shows there is a very sharp transition as temperature is raised, even in relatively short specimen of DNA, leading to dissociation. For some time this has been a serious obstacle in the attempt to model DNA as a one-dimensional system, since one would expect that there is no phase transition for one-dimensional chains [22,24].

However, the theorem on the absence of phase transitions in one dimension only apply to Ising-like (spin) systems, while the situation is far more delicate for systems the state of whose components is described by a continuous local order parameter. Basically, in the former case one describes the system by means of a transfer matrix [22,24] (which is a Markov one, so relevant restriction on its eigenvalues exist – in particular, the fundamental
state cannot be degenerate, and so phase transitions are not possible), while in the latter one one has to use a transfer operator [21]; it is by making use of this formalism, following work done by J.A. Krumhansl and J.R. Schrieffer in 1975, that A. Bishop, Th. Dauxois, and M. Peyrard proved the existence of a “dissociation” phase transition in DNA considered as a one dimensional system [4,5]; see also [6,30].

I do not want to discuss their work here – among other reasons, because I could only provide a bad version of their excellent papers – but I want to stress that their success was not only of mathematical or qualitative nature: their theory compares successfully to experimental data on the detailed (spatiotemporal) dynamics of DNA melting; I want to stress again that they are able to predict not only average quantities, as it should anyway be the case with a Statistical Mechanics approach, but a spatiotemporal pattern: this is a much more severe test of a model, and the model of Peyrard and Bishop [29] for DNA melting which was used (with some minor modifications) in this study did brilliantly pass the test.

Now, what are the characteristics of this model? Well, it is pretty much similar to the models we consider for transcription, and in particular to the Yakushevich one. Indeed it models DNA as a one-dimensional chain, and by singling out one degree of freedom per base – corresponding to “radial” displacements along the axis joining the two bases of a pair – that is, the degree of freedom thought to be the most relevant for the process under study.

Thus the success of Bishop, Peyrard and Dauxois in the study of DNA melting is a witness to the fact that these – apparently, hopelessly oversimplified – models can indeed provide a good understanding, qualitative but also quantitative, of the behaviour of DNA in biologically relevant processes. Simple models can be relevant even in very complicated systems!

I would take this to the extreme, and say that the simplest the model, the greater its value: not only because a simple model can be studied more thoroughly, but first and foremost because if a simple model works, it means it has focused on the right point, i.e. that we have reached an understanding of what are the most relevant aspects (degrees of freedom) of our system for the process under study, and thus it adds more to our knowledge.

Now, comforted by the success of Bishop, Peyrard and Dauxois, we can look back at the simple counting of degrees of freedom in DNA with a more optimistic attitude: it is true that DNA is terribly complicated, but it is also true that it performs a huge variety of essential tasks. While the amazement for how seemingly inconciliable characteristics needed to perform these different tasks can convive in a single molecule should remain, it is possible to think of a process-oriented modelling of DNA, i.e. to obtain a model of DNA which does not have any ambition – or hope – to describe it in general, but only limitedly to a specific (although relevant) process we want to study.

This of course is dependent upon our ability to correctly identify the specific aspects (or degrees of freedom) of DNA which are essential for the process we study. This is the problem whenever we try to formulate a model, and the Peyrard-Bishop model shows that it is possible to do this even in DNA.

A less extreme reduction – keeping a few degrees of freedom per base rather than only one – has been recently considered by Zhang and Collins [46], again obtaining quite
encouraging results. In particular, they compared results obtained by a simplified models with those obtained by an all-atoms simulations, showing that modelling DNA dynamics in terms of a few degrees of freedom is by all means satisfactory, at least for the analysis of specific processes.

Thus, let us come back to discuss what a model of DNA nonlinear excitations which could play a role in transcription should contain.

A first obvious but relevant observation is that the DNA molecule, like any other one, obeys Quantum Mechanics (it should be recalled that both the Peyrard-Bishop and the Yakushevich model are classical); moreover, we are interested in its behaviour in living condition, i.e. in particular at a quite precise temperature (around 300 Kelvin, or a bit more for humans).

Now it happens that at this temperature most of the degrees of freedom of the atoms constituting the DNA molecule will be essentially frozen due to Quantum Mechanics (their excitation energy being much higher), and thus many bonds between atoms can be considered as rigid; on the other side, there are degrees of freedom whose excitation energy is much lower than the thermal scale, so that they can be considered to behave classically.

The consequence of this on modelling is clear: we can safely consider the degrees of freedom of the first kind as non-existent, and concentrate on the remaining – effective – degrees of freedom. These should still be treated quantistically, but for some of them – those of the second kind – we can also use a classical modelling.

It should be clear that the (quite general) reduction to “effective” degrees of freedom is by no means sufficient if we want to consider only one – or even a few – degree of freedom per base: we should have some physical understanding of the process we want to model in order to understand which are the relevant degrees of freedom, which cannot always be just a few.

In the case of transcription the relevant process undergone by the conformation of DNA is its “unwinding” [17,28,33] so that the bases, which usually are stored in the interior of the molecule, as protected as possible from external agents, come to be accessible for reading by the RNAP.

Among the degrees of freedom which are relevant to such a conformational change, there is one which is “softer” than any other, corresponding *grosso modo* to rotations of the bases in a plane orthogonal to the axis of the double helix. Actually such a rotation is, obviously, more complicate than this, but we will investigate the hypothesis that the comparatively small movements around this main rotation are not of fundamental importance (see [17,33]).

It is also reasonable, looking at the geometry of the molecule, to model the conformational change only in terms of this; obviously a lot of other degrees of freedom are involved in such a conformational change, but the idea is that they will play the role of “slave modes”, i.e. there will be small adjustments involving these but which essentially follow the relevant changes undergone by the “master modes”, those of the fundamental degree of freedom.

Such a way of thinking, in terms of relevant variables and slave ones, has been successfull in a wealth of situations, and in Statistical Mechanics [22,24] or in Hydrodynamics [25] one has a very precise way of defining them; however, there is no *a priori* argument
showing that this should be the right way of looking at this specific problem of DNA transcription: we can only justify it \textit{a posteriori} if the model works properly.

On the other side, \textit{if} the model works, this means not only that we have the correct values for coupling constants and so on, but also something more fundamental: that we have individuated the most relevant degree of freedom for the process we are studying.

3. The model and its prediction.

I want now to briefly present the model which has been proposed by L.V. Yakushevich (with some small modification), and its predictions concerning the existence of solitons and their characteristics.

The discussion of the limitations, or of how acceptable are a number of approximations it embodies, will be postponed to the following section.

As I mentioned in the Introduction, several other models have been proposed; I will mention here [2,23,26,27,31,37,38,43,44] (and apologize to other authors I have inadvertently forgotten); in many cases these give predictions similar to those of the Yakushevich model, in particular for the successful ones, but my goal here is not to provide a review of the different models available, and I prefer to restrict my discussion to the model I am familiar with. This is also justified by the fact that this model has been quite successful; and I do not think one can obtain much more without introducing more detailed models and designing specific experiments to test their (presumably, only numerical) predictions. If we look at the Yakushevich “classification” of DNA models [41,42], I would say that her model provides what is reasonable to ask to a model of its level, and that one should now pass to a deeper level to really test the theory and discriminate among different versions.

In the Yakushevich model, we consider rotations of the bases in a plane orthogonal to the double helix axis (which of course is not completely true), and any other movement is not considered. Notwithstanding the fact that the forces involved in hydrogen bonds are highly directional, the forces among the bases are considered to be linear in the distance \(d\) between the extremities interacting to form the H bonds. This will produce a nonlinear dependence on the rotation angles.

An even more severe approximation is introduced concerning the characteristics of the bases and their interactions: all bases are considered as identical, and so the same also holds for base-pair interactions.

Let us consider the interaction among bases on the two chains at site \(n\). If \(r\) is the distance from the center of rotation to the site responsible for the intra-pair interaction, \(\theta_1(n)\) and \(\theta_2(n)\) the angles (both measured in, say, counterclockwise direction [14]) of rotation, and \(L\) the distance at rest, we would then have for the distance \(d\),

\[
d^2 = \left[2L + 2r - r(\cos \theta_1 + \cos \theta_2)\right]^2 + \left[r(\sin \theta_1 + \sin \theta_2)\right]^2.
\]

However, in the Yakushevich model one makes the approximation \(L = 0\) (see next section).

Thus the potential “rotation” energy will be \(V_r = (K_r/2)d^2\), with \(K_r\) a real constant and \(d^2\) the simplified expression corresponding to taking \(L = 0\) in (1). Notice that the force this produces is nonlinear in \(\theta_1, \theta_2\).
Together with this, we will have an elastic interaction with nearest neighbour bases along the two chains, with interaction “torsional” energy \( V_t = (K_t/2)(\theta_1(n) - \theta_1(n - 1))^2 \) and the like for the other chain and for the interaction with bases at site \((n + 1)\).

There is another kind of interaction which should be taken into account[3,12]: bases which are half-pitch of the helix apart are actually near enough in ambient space, and they interact through the formation of water filament; the energy associated to this “helicoidal” interaction is \( V_h = (K_h/2)[\theta_1(n + p) - \theta_2(n)]^2 \) and the like for the other chain, interchanging indices 1 and 2, and for interaction with bases at site \((n - p)\). Here \(p\) is the half-pitch of the helix measured in units of base sites, which for DNA means \(p = 5\). This interaction, although weaker than the other ones, produces a qualitative change in the dispersion relations and in the predictions of the model.

Thus, introducing also the kinetic energy \( K = I/2\dot{\theta}_1(n) + I/2\dot{\theta}_2(n) \), we can finally write the total energy for the (Yakushevich model of the) DNA double chain:

\[
H = \sum_n \frac{1}{2} I \left[ \dot{\theta}_1^2(n) + \dot{\theta}_2^2(n) \right] + \\
+ \sum_n \frac{1}{2} K_t \left[ (\theta_1(n) - \theta_1(n - 1))^2 + (\theta_2(n) - \theta_2(n - 1))^2 \right] + \\
+ \sum_n \frac{1}{2} K_r \left[ 6r^2 + 2r^2 \cos(\theta_1(n) - \theta_2(n)) - 4r^2(\cos(\theta_1(n)) + \cos(\theta_2(n))) \right] + \\
+ \sum_n \frac{1}{2} K_t \left[ (\theta_1(n) - \theta_2(n - p))^2 + (\theta_2(n) - \theta_1(n - p))^2 \right].
\]

From this we can easily derive the equations of motion for the system; these are more usefully written in terms of the variables

\[
\psi_n := \frac{\theta_1(n) + \theta_2(n)}{2}, \quad \chi := \frac{\theta_1(n) - \theta_2(n)}{2}
\]

and are

\[
\ddot{\psi}_n = -\alpha \sin \psi_n \cos \chi_n + \beta (\psi_{n+1} - 2\psi_n + \psi_{n-1}) + \gamma (\psi_{n+p} - 2\psi_n + \psi_{n-p}) \\
\ddot{\chi}_n = -\alpha \sin \chi_n (\cos \psi_n - \cos \chi_n) + \beta (\chi_{n+1} - 2\chi_n + \chi_{n-1}) - \gamma (\chi_{n+p} + 2\chi_n + \chi_{n-p})
\]

(notice the sign differences in the \(\gamma\) terms), where \(\alpha = K_r/I, \beta = K_t/I\) and \(\gamma = K_h/I\).

In these equations we find the different elastic constants \(K_r, K_t, K_h\) or equivalently \(\alpha, \beta, \gamma\); these could be seen as free parameters to be fitted, but it is also possible to give an estimate on their values based on first principles, i.e. on an estimate of the strength of the interactions they should model. The values given in [14,17] are as follows:

\[
K_r = 0.13 \text{ eV}/\text{rad}^2, \quad K_t = 0.025 \text{ eV}/\text{rad}^2, \quad K_h = 0.009 \text{ eV}/\text{rad}^2, \quad I = 3 \cdot 10^{-37} \text{ cm}^2 \text{ g}.
\]

Now, in the analysis of the Yakushevich model it has been usual to pass to the continuum approximation, i.e. having fields \(\psi(x, t), \chi(x, t)\) instead of the sequences of variables \(\{\psi_n(t)\}\) and \(\{\chi_n(t)\}\); in this way the chains of ODEs (4) are replaced by coupled PDEs

\[
\psi_{tt} = -\alpha \sin \psi \cos \chi + \beta \delta^2 \psi_{xx} + \gamma W_+(\psi) \\
\chi_{tt} = -\alpha \sin \chi (\cos \psi - \cos \chi) + \beta \delta^2 \chi_{xx} - \gamma W_-(\chi)
\]

(6)
where $\delta$ represents the distance between consecutive bases along the double chain axis ($\delta \simeq 3.4$ Angstroms), and the nonlinear terms $W_\pm$ are defined by

$$W_\pm(\phi) := \phi(x + p\delta, t) \mp 2\phi(x, t) + \phi(x - p\delta, t).$$

(7)

This passage to the continuum is questionable, and indeed in their study of the melting transition Peyrard and Dauxois have pointed out interesting effects due precisely to the discrete nature of the model (there is a Peierls barrier restraining energy flow); we will discuss this point in next section as well.

We can then analyze (6); first of all – considering the linear terms – we obtain the dispersion relations for the $\psi$ and the $\chi$ branch; these are respectively

$$\omega^2_\psi = \alpha + \beta\delta^2 q^2 + 4\gamma \sin^2(p\delta q/2)$$

$$\omega^2_\chi = \beta\delta^2 q^2 + 4\gamma \cos^2(p\delta q/2).$$

(8)

Notice that now (due indeed to the introduction of the “helicoidal” interactions) the minimum of the dispersion curve, and thus the threshold for excitation of linear modes, corresponds in terms of the natural wavenumber unit $\xi = p\delta q/2$, to $\xi = 1.4$; the corresponding wavelength will be the first to be excited and should act as a seed for the formation of nonlinear structure. Thus, it gives a rough estimate for the size of excitations, which is $\lambda \simeq 11\delta$.

Similarly, the corresponding $\omega = \omega(\xi_0)$ gives a rough estimate for the timescale of small oscillations, which is of $57$ ps, and similarly one obtains an estimate of the amplitude of “linear” oscillations, which is $0.77\pi$. This seems sufficient to ignite the unwinding and the nonlinear regime.

We stress that the minima of the dispersion relations are essentially dependent on the helicoidal interaction term (without this we actually would have phonons in the model). Data for linearized dynamics are in a way even more significant when we analyze DNA melting rather than transcription, but this lies outside our scope here; see [14,17] for a discussion. Also, the presence of helicoidal interactions lead to crossings between the two branches of the dispersion relations, which could play a role in energy transfers [15].

It is remarkable that the estimate of the size of collective motions is of the same order of magnitude as experimental data on the size of transcription bubbles. Again, for melting models the agreement is significant also for other relevant quantities, again providing the right order of magnitude.

Let us now look more precisely to solitons, or solitary waves, in (6). These will be solution of the form $\phi(x, t) = \phi(x - vt) \equiv \phi(z)$, where $\phi = \psi, \chi$. Moreover they should satisfy

$$\lim_{z \to \pm \infty} \phi_z = 0, \quad \lim_{z \to \pm \infty} \phi = 2m\pi$$

(9)

in order to have a finite energy. They can thus be characterized in terms of their winding numbers $\mu = (m_+ - m_-)$ for $\psi$ and $\chi$ [13,14,17]; these corresponds to the phase shift of the solitons.

The lowest energy solitons will be those with $\mu = (1,0)$ and $\mu = (0,1)$, and we concentrate on these.
If we set the helicoidal interaction to zero, it is possible to determine exact solitary wave solutions. These are given by

$$\psi(z) = \arccos\left(\frac{\sinh^2(az) - 1}{\sinh^2(az) + 1}\right)$$

and $\chi(z) = 0$ in the case of the $(1, 0)$ soliton, while the $(0, 1)$ soliton is given by $\psi(z) = 0$ and

$$\chi(z) = \arccos\left(\frac{a^2z^2 - 1}{a^2z^2 + 1}\right).$$

Here $a = (2\alpha / (\beta \delta^2 - v))^{1/2}$, where $v$ is the soliton’s speed.

A numerical integration [13] shows that the introduction of the (weak) helicoidal terms does not really alter the fully nonlinear structures.

The solitonic solution can exist, in principle, with any speed $v < \beta \delta^2$. Moreover, the dependence of their energy on the speed [13,17] is given by a relation of the type $E = c_1/(c_2 - v^2)$, as it should for waves, so that up to high speed the energy is very little sensitive to speed; see [13] for details. This means that in this model there is no selection of soliton speed. A refinement of the model [16] seems to be able to provide such a selection based on some conditions at interface of A-T and G-C regions, but in this model the soliton can not actually exist in proper terms, i.e. as solitary waves travelling along the chain with no dispersion.

As for the size of the soliton, this will correspond to the size $s$ of the region in which $\psi$ differs from its asymptotic value by more than a given quantity $\varepsilon$, and thus will depend on this. We call $z_+, z_-$ the values of $z$ at which $\psi(z_+) = \varepsilon$, $\psi(z_-) = 2\pi - \varepsilon$. For $az \gg 1$, the argument of the arccos in (10) is $f(z) \simeq 1 - 2 \exp[-2aaz]$, and thus if $\cos(\varepsilon) = 1 - \lambda(\varepsilon)$, we have to solve $\lambda(\varepsilon) = 2 \exp[-2aaz] \pm 1$; obviously $z_- = -z_+$, and so $s = |a^{-1} \ln[\lambda(\varepsilon)/2]|$.

As for $a$, when $v = 0$ we get $a = \sqrt{2\alpha / (\beta \delta^2)} := a_0 \delta^{-1}$, and the soliton size measured in base sites units is thus

$$s = \left|\frac{1}{a_0} \ln[\lambda(\varepsilon)/2]\right| \delta.$$  

If we choose $\varepsilon = 5^\circ \simeq 0.044$, we get $\lambda(\varepsilon) \simeq 2.9 \cdot 10^{-7}$ and $\ln[\lambda(\varepsilon)/2] \simeq 15.75$; with the values given in (5) we get $a_0 \simeq 3.22$ and thus it results $s \simeq 5\delta$.

This is smaller than the observed size of the transcription bubble, which is of order 15 - 20 bases, but again we have got the right order of magnitude.

### 4. Limitations of the model.

In the previous section we have seen that the Yakushevich model, with the introduction of “helicoidal” terms, is able to give a prediction of the order of magnitude of several physical quantities for coherent structures and excitations in the DNA chain. This is specially significative when we consider the crude approximations which have been introduced in the formulation of the model, so that we are encouraged to think that the model does
indeed correctly identify the essential degrees of freedom for the dynamics related to the transcription process.

In this section we want instead to emphasize the limitations and the shortcomings of the model; in most cases this will actually be not specific to the Yakushevich model, but common to the whole set of simple models we have advocated above.

We will actually first of all mention a shortcoming of teh Yakushevich model itself, which was noticed by Gonzalez and Martin Landrove [18]: the seemingly unoffensive approximation $L = 0$, see eq.(1), is actually selecting a special case: any nonzero value of $L$ would cause qualitative (and not only quantitative) differences in some basic behaviour of the model; in particular the pairing interaction would be through nonlinear terms only, with no linear ones. As this situation is unphysical, we see that the apparently special choice $L = 0$ does actually produce a “generic” behaviour; this has also suggested to consider the Yakushevich model as an effective one.

Having mentioned this fact, we can pass to aspects which are not really specific of the Yakushevich model, although we will refer to it for the sake of definiteness.

A first obvious limitation is in the fact that bases are considered as identical. Needless to say, this is not the case. If one could be tempted to think that distinguishing between Purines and Pyrimidines only could be acceptable, as a first approximation, in view of the modellization adopted (the physical quantities entering in the model are homogeneous enough for the two purines and for the two pyrimidines, see [16]), considering all bases as equivalent is highly questionable: e.g., masses range from 110 to 150 atomic mass units, and inertia moment from 1500 to 2500 m.u. Angstrom.

Also, the energies involved in the base-base interactions are quite different for A-T and C-G pairs: the H-bond pairing energy is of 7.0 kcal/mole in the first case and 16.8 kcal/mole in the second one; the opening energy is respectively of 4.0 and 7.5 kcal/mole, and the stacking energies in the case of a homogeneous sequence is of 5.4 kcal/mole for a sequence of A-T (or T-A) pairs, and of 8.3 kcal/mole for G-C (or C-G) pairs.

This calls immediately for a more realistic modellization of the DNA chain, taking into account the different characteristics of different bases. It is quite clear, however, that renouncing to the spatial homogeneities of the chain (and to any kind of spatial periodicity as well if we want to model realistic sequences) means that such more realistic models will not have proper solitary waves, i.e. solutions of the form $\phi(x, t) = \phi(x - vt)$ or the discrete analogue.

Naturally, from the point of view of biological significance of solutions we do not need that these are exactly solitary waves propagating with no dispersion and/or change in shape: it is sufficient that they are sufficiently stable over sufficiently long space and time intervals.

Obviously an analytical investigation of a realistic model with a true base sequence is outside reach, so this should be studied numerically. As far as I know, such a numerical study has never been conducted – opposite to the detailed studies conducted by Dauxois and Peyrrard for the related modelling of the dissociation process by a realistic model based on the Peyrard-Bishop one – and so we cannot draw any conclusion in this respect (I do not know of any apriori obstacle to such a numerical study; in particular, it seems that for the nonlinear structure the weak helicoidal interaction, which was essential at the level
of dispersion relations, has very little influence, so that one could study a simple “planar” model [17]).

Partial informations can be extracted for special kind of base sequences (at least in the purine-pyrimidine approximation); thus, for example, alternating regions such as “TATA boxes” would select a zero soliton speed [16]. This is an interesting result, as TATA boxes are indeed know to code the pausing sites for the transcription process.

However, for more general sequences, one has reasons to think that a more detailed Yakushevich-like model could actually be worse than the original simple one from the point of view of correspondence to experimental observations; this could be due to the fact that the structures we are interested in involve several base pairs, so that there could be a mechanism of “effective self-averaging” in the dynamics [16].

Another relevant limitation of Yakushevich-like models is that there seem to be no selection of the soliton speed (see however below): solitons can exist at all speed below the maximal one $v_{\text{max}} = \beta \delta^2$, and energy considerations will not lead to a real selection among low speeds. This is not really encouraging, as transcription speed is not enormously fluctuating: we would expect there is some mechanism to regulate the soliton speed. This would also be essential to the functional role which the soliton is supposed to have, as it is necessary in this picture that the soliton speed is compatible with the speed of the transcription by the RNAP.

One should also recall that, although most of the investigation of the Yakushevich model have been based on the continuum approximation, such an approximation is not justified: actually, as mentioned above, in the case of DNA denaturation Dauxois and Peyrard have pointed out the essential role played by discreteness of the chain.

Again, it seems that no thorough investigation of the Yakushevich model on a chain (even an homogeneous one) have been conducted, so that properly speaking we cannot be sure about the existence and characteristic of soliton-like excitations in this case.

Another relevant point to be recalled is that a soliton in a continuous homogeneous medium would move with zero energy barrer. Essentially, this can be seen realizing that the field Lagrangian will involve $\phi(t)$ and $\phi(x)$, and when $\phi = \phi(x - vt)$ these two terms become $\phi_x^2$ and $v^2 \phi_x^2$: thus a soliton field configuration with zero speed will have an energy $E_0$, and one with a speed $v = \varepsilon$, an energy $E' = (1 + \varepsilon^2)E_0$.

However, the discrete structure of the chain lead to an effective barrer (known as Peierls barrer) for the motion. The result is that excitations of sufficiently low energy can be pinned on-site precisely by the presence of this barrer and thus by the discrete structure. This suggests that in a discrete Yakushevich model only solitons of sufficient high energy and thus speed could actually move along the chain, passing the Peierls barrer.

It should be noticed that this could provide the required mechanism for soliton speed, imposing $v > v_0$; the significance of this would of course depend on the actual value of $v_0$ and how it compares with the observed speed of the transcription bubble.

Another major limitation of the Yakushevich – and similar – models is precisely the fact of focusing on the rotational motion connected with the reading of the DNA sequence by RNAP: indeed, the rigidity (or softness) of the concerned degree of freedom is quite comparable to the one of “radial” motion as considered in the Peyrard-Bishop model; thus, it would be highly desirable to consider these two degrees of freedom together, and combine
the Peyrard-Bishop and the Yakushevich models into a single one.

Finally, the models – such as Yakushevich’s one – studied in connection with the problem of DNA transcription are all purely deterministic and only concern the DNA chain itself, i.e. with no interaction with its environment. One hopes that in a large part the chemico-physical parameters describing the environment could somehow be embodied in effective values of the constants defining the model (or one could limit to consider “standard” environment), but however the DNA has to work at finite, actually quite different from zero, temperature: thus unavoidably one has to consider thermal energy of the environment and energy exchanges involving this.

In considering this problem, one could resort to Statistical Mechanics considerations (which lie outside the scope of the present short report); or more in the spirit of Yakushevich-like models one should introduce random forces modelling thermal effects. It is not clear, nor – as far as I am aware – it has been studied if the solitons would survive the introduction of such random forces.

On the other side, the role of thermal energy is not necessarily negative from the point of view of existence of solitons: it is well possible that these same random forces would help the initial formation of nonlinear structures by pumping energy into the more easily excitable modes until they ignite the formation of soliton-like collective structures.

It is maybe worth mentioning that statistical approaches to inhomogeneous models have been also considered recently; I will just mention the work of Hisakado and Wadati [19], where the distribution of masses and physical constants of the bases is a gaussian white noise, and the more specific work of Homma [20], which sets the basis for a statistical mechanics treatment of double chain models with sine-like nonlinearities, i.e. of the kind we consider here.

In this respect, it should be recalled that the whole process of transcription is strongly dependent on temperature (as is also that of denaturation), and in in vitro experiments where temperature is varied the transcription bubble forms sharply in a very small temperature range, see [17] and references therein: thus we are compelled to introduce dependence upon temperature in any modellization of the process.

This problem is considered e.g. in [36]; see also [45] for consideration of the environment in the somehow related problem of a single polyethilene chain.

5. Summary and conclusions.

I have given a short discussion of the idea that nonlinear excitations could play a role in the process of DNA transcription, i.e. that the transcription bubble could correspond to a solitary wave travelling along the chain, which the RNAP could then “surf” in order to access the base sequence with no energy to provide for opening the double helix.

I have discussed at some length the general idea of providing a simple model for a specific DNA process, and argued that despite the tremendous complexity of the DNA model this approach is not bound to fail. I have then recalled the main features of the model proposed by Yakushevich, mentioning some encouraging achievements and several limitations.
These limitations, however, more than being inherent to the model are limitation of the studies conducted so far. It is clear that the model is too simple to be valid as it is, and that it is needed to go “one step further” in the Yakushevich classification of DNA models [41,42], but only a more thorough analysis can focus on the detailed refinements which are needed.

In particular, I have pointed out several directions in which I believe it is necessary to generalize the model and to investigate its behaviour, such as considering real base sequences and thermal effects.

I hope this conference can, among other things, also stimulate a reflection on this theme, and maybe new works to analyze more realistic models of the nonlinear dynamics involved in DNA transcription.

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