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# Traveling Waves and Impact Parameter Correlations in QCD Beyond the 1D Approximation 

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# Traveling Waves and Impact Parameter Correlations in QCD Beyond the 1D Approximation 

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## Dissertation

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Dedicated to my parents, who planted the germ of everything I have accomplished.

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# Traveling Waves and Impact Parameter Correlations in QCD Beyond the 1D Approximation 

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The theory of quantum chromodynamics (QCD) predicts that at high energies, such as those investigated in deep inelastic scattering experiments, hadrons evolve into dense gluonic states described by the BFKL equation, and at very high densities, the more general BK equation. In certain approximations, the BK equation reduces to a well studied reaction-diffusion type nonlinear partial differential equation, the FKPP equation, for which analytical results are known. In this work, we model the BK equation using a classical branching process rooted in the dipole model of QCD evolution. Because the BK equation is inherently two dimensional, our model allows dipole impact parameters to occupy the full transverse space. A one dimensional limit of this model is studied as well. Results are compared with the predictions of the FKPP equation, and correlations between evolution at different impact parameters are presented. The general features of previously studied one dimensional impact parameter models are verified, but the details are refined in what we believe to be a more accurate model.

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## Part I

## Background

## 1 Introduction

Much effort has been applied to the understanding of a hadron's transition from a dilute parton gas to a saturated CGC (Color Glass Condensate). While the DGLAP (Dokshitzer, Gribov, Lipatov, Altarelli, Parisi) equation could explain data collected at DESY-HERA at very large $Q^{2}$, the investigation of the scaling region at moderate $Q^{2}$ and very small $x \sim Q^{2} /\left(Q^{2}+s\right)$ prompted the application of the integro-differential BFKL (Balitsky, Fadin, Kuraev, Lipatov) equation, which resums infrared logarithms $(\log 1 / x)$. [?]
In its original formulation, the BFKL equation can be derived from the infinite sum of ladder diagrams of Reggeized gluons in the t-channel, as described in [6]. This derivation is known as the "BFKL pomeron" or "hard pomeron", giving the Regge trajectory $\alpha_{P}(t)=1+4 \bar{\alpha}_{s} \ln 2$. However, in the mid 90s, Mueller was able to rederive the BFKL equation in a much simpler s-channel picture and show that the BFKL pomeron is equivalent to a formulation describing dipole splittings in transverse space[11, 12]. A set of color dipoles comprise a so-called onium configuration, in which the emission of new gluons gives rise to new dipoles. Evolution consists of "parent" dipoles splitting into "daughter" dipoles with a characteristic probability

$$
\begin{equation*}
\frac{d P_{x_{01} \rightarrow x_{02}, x_{12}}}{d Y}=\frac{x_{01}^{2} d^{2} \mathbf{x}_{2}}{x_{02}^{2} x_{12}^{2}} \tag{1}
\end{equation*}
$$

The amplitude of a photonic probe interacting with such a highly evolved hadron is roughly proportional to the number of dipoles in the hadron having the same approximate impact parameter and size as the $q \bar{q}$ dipole into which the probe splits.

In the context of the dipole model, the BK (Balitsky, Kovchegov) equation-essentially the BFKL equation modified by a nonlinear term responsible for saturation in the CGC regime-has been studied in a variety of analytical and computational ways in the past decade. The full BK equation in transverse space reads


Figure 1: Phase diagram of a hadron in deep inelastic scattering

$$
\begin{equation*}
\frac{\partial N\left(\mathbf{x}_{01}, Y\right)}{\partial Y}=\frac{\bar{\alpha}}{2 \pi} \int_{\rho} d^{2} \mathbf{x}_{2} \frac{x_{01}^{2}}{x_{02}^{2} x_{12}^{2}} 2 N\left(\mathbf{x}_{02}, Y\right)-N\left(\mathbf{x}_{01}, Y\right)-N\left(\mathbf{x}_{02}, Y\right) N\left(\mathbf{x}_{12}, Y\right) \tag{2}
\end{equation*}
$$

The first two terms on the right hand side represent the increase in the amplitude due to branching diffusion, the third term a virtual correction necessary to normalize the onium wavefunction [17], and the final term the nonlinearity that restores unitarity to the BFKL equation.

One of the most exciting theoretical developments of the past decade has been the discovery that for fixed impact parameter collisions, the BK equation belongs to the universality class of the FKPP (Fisher Kolmogorov, Petrovsky, Piscounov) equation[14, 15]. That is, an analogy was noted between high energy QCD evolution and a well studied reaction-diffusion equation. With the appropriate transformations, the scattering amplitude can be put into the form

$$
\begin{equation*}
\partial_{t} u(t, x)=\partial_{x}^{2} u(t, x)+u(t, x)-u^{2}(t, x) \tag{3}
\end{equation*}
$$

the solution of which describes a traveling wave. The time, $t$, is analogous to the rapidity, $Y$, and spatial coordinate $x$ to the dipole momentum. It is thereby possible to speak of a saturation wave front, $\rho_{s}$, that travels to smaller dipole sizes as collision
energy increases.


Figure 2: Traveling wave solution to the FKPP equation [2]

An important caveat to the application of the FKPP equation is that it is a mean field limit of the true stochastic evolution equations. Due to the discrete nature of an onium state consisting of a finite number of dipoles, fluctuations in dipole number must play a role in the evolution. Because the true stochastic equations are not known and their formulation would probably require a more sophisticated understanding of the saturation mechanism than is presently available, many researchers have taken to monte carlo computer modeling of stochastic splittings. This continues to be a very active field of research [20, 21, 22, 23, 24, 25]. Two of the most recent of these in particular [24, 25] have informed the study described in this manuscript. It will be explained what has been accomplished so far and how it can be extended using a full two dimensional model.

## 2 History of the BFKL Equation

### 2.1 Regge Theory and the origins of the Pomeron

Before the advent of QCD, a variety of other approaches were used to study strong interactions, some of which are still useful today. Regge theory, a branch of S-matrix theory, was for instance successfully used to predict the rise of hadronic cross-sections at small $x$, or increasing center of mass energy. During the sixties when the fundamentals of strong interactions were not yet known, studies focused on the exchange of massive mesons, as in the Yukawa theory of nuclear force. At that time it was postulated (by Chew and Frautschi [3, 4], for example) that there were no elementary strongly interacting particles among hadrons, i.e. mesons and baryons, as it appeared as a consequence of Regge Theory that all hadrons are bound states or resonances with interlocking angular momentum states. To this end a substantial attempt was made to explain all of strong interactions through studying the implications of a number of assumptions about the S-matrix. The argument was that if the strongly interacting particles that were known obeyed a self-consistent theory of the S-matrix, then the need for elementary particles of the strong force would be obviated, yielding a "bootstrap" theory, as it was called.

It was not until detailed data of the nucleon structure functions was obtained from inelastic electron-proton scattering at Stanford Linear Accelerator in 1969 that the physics community came to accept the existence of spin $\frac{1}{2}$ "partons", as Feynman dubbed them, which comprise the nucleon. Although this marked the shift toward what was the beginning of QCD (and the decreasing popularity of the S-matrix approach, especially with regards to phenomenology), it is worth reflecting of the substantial successes of S-matrix theory and how they have shed light on much later developments in QCD. Some insights from S-matrix theory still await a proper QCD treatment while others lie beyond the reach of a perturbative theory like QCD.

We will now give an abbreviated tour of Regge theory, in which amplitudes of strong interaction processes are expanded in terms of partial waves:

$$
\begin{equation*}
A_{a \bar{c} \rightarrow \bar{b} d}(s, t)=\sum_{l=0}^{\infty}(2 l+1) a_{l}(s) P_{l}(1+2 t / s) \tag{4}
\end{equation*}
$$

or by crossing symmetry,


Figure 3: The Chew Frautschi plot of mesons' mass squared versus spin. [7]

$$
\begin{equation*}
A_{a b \rightarrow c d}(s, t)=\sum_{l=0}^{\infty}(2 l+1) a_{l}(t) P_{l}(1+2 s / t) \tag{5}
\end{equation*}
$$

where $P_{l}(z)$ are Legendre polynomials and $a_{l}(s)$ are called a partial wave amplitudes. (5) can be rewritten as a contour integral in the complex angular momentum plane in what is known as a Sommerfeld-Watson transform. The contour surrounds the positive $x$-axis so that the residues reproduce the sum in (5):

$$
\begin{equation*}
A(s, t)=\frac{1}{2 i} \oint_{C} d l(2 l+1) \frac{a(l, t)}{\sin \pi l} P(l, 1+2 s / t) \tag{6}
\end{equation*}
$$

$a(l, t)$ and $P(l, 1+2 s / t)$ are analytic continuations of the functions in (5). If we consider the Regge region $s \gg|t|$, we can expand $P_{l}(z)$ as

$$
\begin{equation*}
P_{l}(1+2 s / t) \xrightarrow{s \gg t} \frac{\Gamma(2 l+1)}{\Gamma^{2}(l+1)}\left(\frac{s}{2 t}\right)^{l} \tag{7}
\end{equation*}
$$

This allows us to conveniently deform the contour in (6) to a vertical line on which $\Re \mathrm{e}(l)<0$, causing $\left(\frac{s}{2 t}\right)^{l}$ to vanish at large $s$. In the process of deforming the contour, however, we pick up poles in the $l$ plane known as Regge poles. The residue of the pole with the largest real part leads to the amplitude behavior

$$
\begin{equation*}
A(s, t) \stackrel{s \rightarrow \infty}{\sim} s^{\alpha(t)} \tag{8}
\end{equation*}
$$

Recalling that $\alpha(t)$ is an angular momentum, one can learn about this function by
plotting low lying mesons with spin $J_{i}$ and mass $m_{i}$, as done on figure 3. It then becomes immediately obvious that $J_{i}=\alpha\left(m_{i}^{2}\right)$ is a linear function, i.e. $\alpha(t)=$ $\alpha(0)+\alpha^{\prime} t$. The intercept of this plot has a special meaning: the optical theorem at large $s$ gives the forward total cross-section as

$$
\begin{equation*}
\sigma_{t o t} \propto s^{\alpha(0)-1} \tag{9}
\end{equation*}
$$

Thus, the Regge intercept determines the total cross section. From figure 3, it appears the intercept is about .5 , implying that the so-called Reggeons in the figure contribute

$$
\begin{equation*}
\sigma_{t o t} \propto s^{-0.5} \tag{10}
\end{equation*}
$$

to the total cross-section. But this is not at all what is observed! Instead, data shows that cross-sections rise starting at $\sqrt{s} \gtrsim 10 \mathrm{GeV}$. In the late 1950s, Pomeranchuk proved that any scattering process in which there is charge exchange exhibits an asymptotically vanishing cross-section. Therefore, there must be a exchange with vacuum quantum numbers that causes the cross-section to rise. This Regge trajectory is called the Pomeron ${ }^{11}$. Later after the advent of QCD, it was conjectured that the integer values of the Pomeron trajectory $\alpha_{\mathbb{P}}(t)$ might correspond to bound states of gluons, or glueballs. Proving the existence of such entities remains one of the great remaining experimental challenges of high energy QCD.

### 2.2 The hard Pomeron attained through QCD ladder diagrams

Once perturbative QCD techniques had become well established, it was naturally wondered whether Pomeron behavior could be derived from pQCD. Copious detail on this program can be found in [6], the results of which we will now briefly touch on. Computing infinite ladder diagrams such as figure 4 left can reproduce the Pomeron behavior of (9). Slashes through vertical gluons indicate they have been "Reggeized", i.e. each is a sum of infinite ladder rungs such that the gluon propagator is replaced by

[^0]

Figure 4: Left: A ladder diagram of Reggeized gluons representing Pomeron exchange. Right: Diagram illustrating the integral equation for the Mellin transformed amplitude, $f(\omega)$.

$$
\begin{equation*}
\tilde{D}_{\mu \nu}\left(s_{i}, k_{i}^{2}\right)=\frac{i g_{\mu \nu}}{\mathbf{k}_{i}^{2}}\left(\frac{s_{i}}{\mathbf{k}^{2}}\right)^{\epsilon\left(k_{i}^{2}\right)} \tag{11}
\end{equation*}
$$

where $i$ stands for the $i$ th rung and $s_{i}=\left(k_{i-1}-k_{i+1}\right)^{2}$ is the squared center of mass energy coming into the $i$ th rung.

One may write an integral equation shown diagrammatically in figure 4 right and solve for the Mellin transformed amplitude at zero momentum transfer, $f\left(\omega, \mathbf{k}_{1}, \mathbf{k}_{2}, \mathbf{0}\right)$, as such: [6]

$$
\begin{equation*}
f\left(\omega, \mathbf{k}_{1}, \mathbf{k}_{2}, \mathbf{0}\right) \approx \frac{1}{\pi k_{1} k_{2}} \int_{-\infty}^{\infty} \frac{d \nu}{2 \pi}\left(\frac{k_{1}^{2}}{k_{2}^{2}}\right)^{i \nu} \frac{1}{\omega-\omega_{0}+a^{2} \nu^{2}} \tag{12}
\end{equation*}
$$

with

$$
\begin{equation*}
\omega_{0}=4 \bar{\alpha}_{s} \ln 2 \tag{13}
\end{equation*}
$$

and $\nu$ the anomalous dimension of the BFKL eigenvalue function, which we will later cover in detail. Performing the contour integration and inverting the Mellin transform,

$$
\begin{equation*}
F\left(s, \mathbf{k}_{1}, \mathbf{k}_{2}, \mathbf{0}\right) \approx \frac{1}{\sqrt{\mathbf{k}_{1}^{2} \mathbf{k}_{2}^{2}}}\left(\frac{s}{k^{2}}\right)^{\omega_{0}} \frac{1}{\sqrt{\pi \ln \left(s / k^{2}\right)}} \frac{1}{2 \pi a} \exp \left(-\frac{\ln ^{2}\left(\mathbf{k}_{1}^{2} / \mathbf{k}_{2}^{2}\right)}{4 a^{2} \ln \left(s / k^{2}\right)}\right) \tag{14}
\end{equation*}
$$

The full $q \bar{q}$ forward elastic scattering amplitude is then

$$
\begin{equation*}
\frac{\mathcal{A}^{(1)}(s, 0)}{s}=4 i \alpha_{s}^{2} \delta_{\lambda_{1}^{\prime} \lambda_{1}} \delta_{\lambda_{2}^{\prime} \lambda_{2}} G_{0}^{(1)} \int \frac{d^{2} \mathbf{k}_{1}}{k_{1}^{2}} \frac{d^{2} \mathbf{k}_{2}}{k_{2}^{2}} F\left(s, \mathbf{k}_{1}, \mathbf{k}_{2}, \mathbf{0}\right) \tag{15}
\end{equation*}
$$

and thus,

$$
\begin{equation*}
\sigma_{t o t} \sim s^{\omega_{0}}=s^{\alpha_{\mathbb{P}}(0)-1}=s^{4 \bar{\alpha}_{s} \ln 2} \tag{16}
\end{equation*}
$$

So we see that the pQCD ladder diagram calculation successfully predicts the Pomeron trajectory required for the rise of the total cross-section.


Figure 5: Inclusive deep inelastic scattering for $e^{-} p \rightarrow e^{-} X$

## 3 Dipole Formulation of BFKL Equation

### 3.1 Description of the dipole model

So far we have looked at the BFKL equation from the standpoint of t -channel interactions of $\gamma^{*} p \rightarrow X$. However, a much simpler method of deriving the BFKL equation was achieved in the s-channel picture by Mueller [11], in which the evolution takes place in the target as one boosts it to greater rapidity. In this approach, the target interacts with the probe as an "onium" state of quantum fluctuations. An onium comprises a high occupancy Fock state when the interaction energy is large. Using the onium wavefunction to calculate the dipole cross-section, other useful deep inelastic scattering observables may be calculated.

The idea for calculating the dipole cross-section had been popular before Mueller used it to rederive the BFKL equation [33][34]. In a process such as $e^{-} p \rightarrow e^{-} X$ (see figure 5), the dominant contribution to the scattering cross-section comes from photon's dissociation into a quark-antiquark color-singlet state that strongly interacts with the proton (see figure 6). This approach is only legitimate when the dissociation time of the photon is large compared to interaction time with the proton. We can estimate these times using energy uncertainty as follows [7]. Let the four-momentum of the photon, quark, and antiquark be, respectively,

$$
\begin{equation*}
q=\left(q_{0}, 0,0, q_{3}\right) \quad k_{1}=\left(E_{1}, \mathbf{k}_{T}, z q_{3}\right) \quad k_{2}=\left(E_{2},-\mathbf{k}_{T},(1-z) q_{3}\right. \tag{17}
\end{equation*}
$$



Figure 6: Photon dissociation into quark-antiquark pair and interaction with hadron. A cut of the total cross-section is displayed.
where $z$ is the fraction of the photon momentum carried by the quark ( $0 \leq z \leq 1$ ), and $\mathbf{k}_{T}$ is the two dimensional transverse momentum of the quark. The dissociation time for the photon is then given by

$$
\begin{equation*}
\tau_{d i s}=\frac{1}{\left|q_{0}-E_{1}-E_{2}\right|} \tag{18}
\end{equation*}
$$

Using expansions in the large $q_{3}$ limit, $E_{1} \approx z q_{3}+\frac{m_{f}^{2}+\mathbf{k}_{T}^{2}}{2 z q_{3}}, E_{2} \approx(1-z) q_{3}+\frac{m_{f}^{2}+\mathbf{k}_{T}^{2}}{2(1-z) q_{3}}$, $q_{0} \approx q_{3}-\frac{Q^{2}}{2 q_{3}}$,

$$
\begin{equation*}
\tau_{d i s} \approx \frac{1}{\left|-\frac{Q^{2}}{2 q_{0}}-\frac{m_{f}^{2}+\mathbf{k}_{T}^{2}}{2 z(1-z) q_{0}}\right|} \tag{19}
\end{equation*}
$$

If we take the interaction time of the dissociated photon with the proton in its rest frame to be of the order of the proton confinement radius $1 / \Lambda$, and set $\left|\mathbf{k}_{T}\right| \approx \Lambda$, our timescale comparison yields

$$
\begin{gathered}
\tau_{d i s} \gg \frac{1}{\Lambda} \\
2 q_{0} m_{p} \gg \frac{m_{p}}{\Lambda}\left(Q^{2}+\frac{m_{f}^{2}+\Lambda^{2}}{z(1-z)}\right)
\end{gathered}
$$



Figure 7: Diffractive deep inelastic scattering

$$
\begin{equation*}
W^{2} \gg \frac{m_{p}}{\Lambda}\left(Q^{2}+\frac{m_{f}^{2}+\Lambda^{2}}{z(1-z)}\right) \tag{20}
\end{equation*}
$$

since $W^{2}=(p+q)^{2}=m_{p}^{2}-Q^{2}+2 m_{p} q_{0}$ in the proton rest frame. 20 tells us that unless $z$ is close to 0 or $1, W^{2} / Q^{2} \gg 1$. This condition has a special significance in deep inelastic scattering-recalling the definition of the Bjorken $x$,

$$
\begin{equation*}
x:=\frac{Q^{2}}{2 p \cdot q}=\frac{Q^{2}}{(p+q)^{2}-m_{p}^{2}-q^{2}} \approx \frac{Q^{2}}{W^{2}+Q^{2}} \tag{21}
\end{equation*}
$$

We see that $W^{2} / Q^{2} \gg 1$ at large energies implies we are in the small $x$ regime. Therefore, for the high energy processes we will be considering, the dipole picture is appropriate. Note that this method differs from the usual deep inelastic picture in which a parton is knocked out by the virtual photon in that the dipole is interacting with the gluonic field of the hadron, as opposed to a single parton.

Deep inelastic scattering experiments, such as HERA, have been among the most fruitful for the application of the dipole model. Deep inelastic scattering itself is good testing ground for high energy QCD since the photon kinematics are contained in the measurement of the outgoing lepton, yielding $Q^{2}$. Models for the dipole cross-section have successfully been applied to inclusive and diffractive events at HERA [35, 36, 37] (see figure 7 for an illustration of the latter).
In order to derive QCD evolution equations, we should focus our attention on the wavefunction of the onium state of the target hadron. This state is built from succes-


Figure 8: Quark-antiquark pair interacting with an evolved target. A cut of the total cross-section is displayed.
sive splittings of the original valence partons of the target until dense gluonic states comprise the target at high energy. This process is called a gluonic cascade, a still shot of which is shown in figure 8. Because quarks or gluons splitting into a gluon exhibit a logarithmic singularity in $z$ [1], soft gluons dominate in the small $x$ limit or alternatively in the large rapidity limit, as $y=\ln 1 / x$. In the limit of large number of colors $\left(N_{c}\right)$, each emitted gluon is treated as a zero-size quark-antiquark pair ${ }^{2}$, as shown in figure 9. Note, however, that the dipoles are of finite size, as can also be seen in the figure. This is a potential source of confusion, as we usually think of a dipole as being the limit of zero separation between a charge and anti-charge, although in this case the color dipoles are finite size.
A major advantage to the dipole-onium interaction model is that the cross-section for the subprocess shown in figure 8 factorizes:

$$
\begin{equation*}
\sigma^{\gamma^{*} p}\left(Y, Q^{2}\right)=\int d^{2} b d^{2} x_{01} \int_{0}^{1} d z\left|\psi_{\gamma^{*}}\left(z, x_{01} Q\right)\right|^{2} \sigma_{\text {dipole }}\left(Y, x_{01}\right) \tag{22}
\end{equation*}
$$

where $\psi_{\gamma^{*}}\left(z, x_{01} Q\right)$ is the photon wavefunction for splitting into a quark-antiquark dipole of size $x_{01}, z$ the longitudinal momentum fraction of the quark, and $\sigma_{\text {dipole }}$ the dipole forward scattering amplitude.

[^1]

Figure 9: Illustration of dipoles in the evolved target from figure 8. Each dipole is indicated by a double-headed arrow.

### 3.2 QCD evolution using color dipoles

### 3.2.1 Single emitted gluon wavefunction

With the dipole model of hadron evolution we can now see how QCD evolution equations, in particular the BFKL equation, can be obtained. We will follow the seminal paper by Mueller [11] with the addition of some omitted details. The accuracy of our calculation will be leading logarithmic such that the $\left(\alpha \ln \frac{1}{z_{0}}\right)^{n}$ contribution to the square of the onium wavefunction will be computed for $n$ soft gluons with momentum between $z_{0} p$ and $p$. Using the usual Feynman rules for a gluon and quark vertex, the diagrams in figure 10 yield the following contribution to the momentum space onium wavefunction:

$$
\begin{equation*}
\psi_{\alpha \beta}^{(1) a}\left(\mathbf{k}_{1}, \mathbf{k}_{2} ; z_{1}, z_{2}\right)=-g T^{a}\left[\psi_{\alpha \beta}^{(0)}\left(\mathbf{k}_{1} ; z_{1}\right)-\psi_{\alpha \beta}^{(0)}\left(\mathbf{k}_{1}+\mathbf{k}_{2} ; z_{1}\right)\right] \frac{\mathbf{k}_{2} \cdot \epsilon_{2}^{\lambda}}{k_{2}^{2}} \tag{23}
\end{equation*}
$$

where $a$ is the color index of the emitted gluon, $T^{a}$ the $\mathrm{SU}(3)$ generator, $\alpha$ and $\beta$ spinor indices, $z_{n}:=k_{n}^{+} / p^{+}$the fractional momentum of the original quark-antiquark pair (in lightcone coordinates), $\epsilon_{2}^{\lambda}$ the polarization vector of the emitted gluon with helicity $\lambda$, and $\psi^{(n)}$ is the wavefunction when $n$ soft gluons have been emitted.

We will now transform the momentum space wavefunction to transverse space where a significant simplification takes place: in the high energy limit the emission of small $z$, or soft, gluons dominates, and the transverse coordinates of the parent partons are not affected by subsequent evolution of the system. Thus, each dipole evolves


Figure 10: Single gluon emission from quark-antiquark pair
independently of the others. Their transverse coordinates are said to be "frozen". Fourier transforming to transverse space,

$$
\begin{equation*}
\psi_{\alpha \beta}^{(1) a}\left(\mathbf{x}_{1}, \mathbf{x}_{2} ; z_{1}, z_{2}\right)=\int \frac{d^{2} \mathbf{k}_{2}}{(2 \pi)^{2}} \int \frac{d^{2} \mathbf{k}_{1}}{(2 \pi)^{2}} e^{i \mathbf{k}_{1} \cdot \mathbf{x}_{1}+i \mathbf{k}_{2} \cdot \mathbf{x}_{2}} \psi_{\alpha \beta}^{(1) a}\left(\mathbf{k}_{1}, \mathbf{k}_{2} ; z_{1}, z_{2}\right) \tag{24}
\end{equation*}
$$

Substituting (23) into (24),

$$
\begin{gather*}
=g T^{a} \int \frac{d^{2} \mathbf{k}_{2}}{(2 \pi)^{2}} e^{i \mathbf{k}_{2} \cdot \mathbf{x}_{2}}\left(\psi_{\alpha \beta}^{(0)}\left(\mathbf{x}_{1} ; z_{1}\right)-\int \frac{d^{2} \mathbf{k}_{1}^{\prime}}{(2 \pi)^{2}} e^{i\left(\mathbf{k}_{1}^{\prime}-\mathbf{k}_{2}\right) \cdot \mathbf{x}_{1}} \psi_{\alpha \beta}^{(0)}\left(\mathbf{k}_{1}^{\prime} ; z_{1}\right)\right) \frac{\mathbf{k}_{2} \cdot \epsilon_{2}^{\lambda}}{k_{2}^{2}} \\
=g T^{a} \psi_{\alpha \beta}^{(0)}\left(\mathbf{x}_{1} ; z_{1}\right) \int \frac{d^{2} \mathbf{k}_{2}}{(2 \pi)^{2}}\left(e^{i \mathbf{k}_{2} \cdot\left(\mathbf{x}_{2}-\mathbf{x}_{0}\right)}-e^{i \mathbf{k}_{2} \cdot\left(\mathbf{x}_{2}-\mathbf{x}_{1}\right)}\right) \frac{\mathbf{k}_{2} \cdot \epsilon_{2}^{\lambda}}{k_{2}^{2}} \tag{25}
\end{gather*}
$$

At this point we will need to prove the following Hankel transform:

$$
\begin{equation*}
\int d^{2} \mathbf{k} e^{i \mathbf{k} \cdot \mathbf{x}} \frac{\mathbf{k} \cdot \epsilon}{k^{2}}=-2 \pi i \frac{\mathbf{x} \cdot \epsilon}{x^{2}} \tag{26}
\end{equation*}
$$

We can demonstrate (26) as follows:

$$
\begin{aligned}
L H S & =\sum_{j=1,2} \int d^{2} \mathbf{k} e^{i \mathbf{k} \cdot \mathbf{x}} \frac{k_{j} \epsilon_{j}}{k^{2}} \\
& =-i \sum_{j=1,2} \frac{\partial}{\partial x_{j}} \int d^{2} \mathbf{k} e^{i \mathbf{k} \cdot \mathbf{x}} \frac{\epsilon_{j}}{k^{2}} \\
& =2 \pi i \sum_{j=1,2} \hat{e}_{j} \cdot \nabla_{x} \int_{0}^{\infty} d k J_{0}(k x) \frac{\epsilon_{j}}{k} \\
& =-2 \pi i \sum_{j=1,2} \hat{e}_{j} \cdot \hat{x} \int_{0}^{\infty} d k J_{1}(k x) \epsilon_{j} \\
& =-2 \pi i \sum_{j=1,2} \frac{\hat{e}_{j} \cdot \hat{x} \epsilon_{j}}{x} \\
& =-2 \pi i \frac{\mathbf{x} \cdot \epsilon}{x^{2}}
\end{aligned}
$$

Using (26) in (25), we obtain

$$
\begin{equation*}
=-\frac{i g T^{a}}{2 \pi} \psi_{\alpha \beta}^{(0)}\left(\mathbf{x}_{1} ; z_{1}\right)\left(\frac{\mathbf{x}_{20}}{x_{20}^{2}}-\frac{\mathbf{x}_{21}}{x_{21}^{2}}\right) \cdot \epsilon_{2}^{\lambda} \tag{27}
\end{equation*}
$$

where a Hankel transform has been performed in the last step. Note that $\mathbf{x}_{0}=0$ in the above, and $\mathbf{x}_{20}:=\mathbf{x}_{2}-\mathbf{x}_{0}, \mathbf{x}_{21}:=\mathbf{x}_{2}-\mathbf{x}_{1}$. Now let us calculate the squared and summed wavefunction. If the squared and summed wavefunction for zero gluons present is

$$
\begin{equation*}
\Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right):=\sum_{\alpha \beta}\left|\psi_{\alpha \beta}^{(0)}\left(\mathbf{x}, z_{1}\right)\right|^{2} \tag{28}
\end{equation*}
$$

then similarly, that for one gluon present is

$$
\Phi^{(1)}\left(\mathbf{x}_{1}, z_{1}\right):=\int d^{2} \mathbf{x}_{2} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \sum_{\alpha \beta} \frac{1}{2} \sum_{\lambda=1,2} \sum_{a}\left|\psi_{\alpha \beta}^{(1) a}\left(\mathbf{x}_{1}, \mathbf{x}_{2} ; z_{1}, z_{2}\right)\right|^{2}
$$

$z_{0}$ serves as a lower cutoff to the emitted gluon momentum, $z_{2}$. The largest momentum the gluon can possess is $z_{1}$ in the leading logarithmic approximation.

$$
\begin{gather*}
=\frac{1}{2} \frac{g^{2}}{(2 \pi)^{2}} \sum_{a} T^{a} T^{a} \int d^{2} \mathbf{x}_{2} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \sum_{\alpha \beta}\left|\psi_{\alpha \beta}^{(0)}\left(\mathbf{x}, z_{1}\right)\right|^{2} \sum_{\lambda=1,2}\left[\left(\frac{\mathbf{x}_{20}}{x_{20}^{2}}-\frac{\mathbf{x}_{21}}{x_{21}^{2}}\right) \cdot \epsilon_{2}^{\lambda}\right]^{2} \\
=\frac{\alpha N_{c}}{\pi} \int \frac{d^{2} \mathbf{x}_{2}}{2 \pi} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right)\left(\frac{\mathbf{x}_{20}}{x_{20}^{2}}-\frac{\mathbf{x}_{21}}{x_{21}^{2}}\right)^{2} \tag{29}
\end{gather*}
$$

where we have used the strong coupling constant $\alpha_{s}=\frac{g^{2}}{4}$, the trace over $\sum_{a} T^{a} T^{a}=$ $N$ in the adjoint representation of $\mathrm{SU}(\mathrm{N})$, and the polarization sum was evaluated with $\epsilon^{1}=(0,1,0,0)$ and $\epsilon^{2}=(0,0,1,0)$. After foiling the term in parenthesis in 29 and some algebraic simplification we arrive at

$$
\begin{equation*}
\Phi^{(1)}\left(\mathbf{x}_{1}, z_{1}\right)=\frac{\alpha N_{c}}{\pi} \int \frac{d^{2} \mathbf{x}_{2}}{(2 \pi)} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) \tag{30}
\end{equation*}
$$

At this point we might want to pause to see what we have gained. Notice that the momentum space representation of single gluon emission,

$$
\begin{equation*}
\Phi^{(1)}\left(\mathbf{k}_{1}, z_{1}\right)=\frac{1}{(2 \pi)^{2}} \int d^{2} \mathbf{k}_{2} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \frac{1}{2} \sum_{\lambda, a, \alpha \beta}\left|\psi_{\alpha \beta}^{(1) a}\left(\mathbf{k}_{1}, \mathbf{k}_{2}, z_{1}, z_{2}\right)\right|^{2} \tag{31}
\end{equation*}
$$

does not exhibit the same clean factorization as (30), which is written as an integral of the zero gluon, bare quark-antiquark wavefunction squared. The simplicity of (30) will allow us to generalize the onium wavefunction to include $n$ soft gluons. Also, we will see the kernel of the spatial integral, $x_{10}^{2} / x_{20}^{2} x_{21}^{2}$ play a significant role later in this manuscript.

### 3.2.2 $n$ emitted gluon wavefunction

For notational simplicity, let us make use of the following Jacobian,

$$
\begin{equation*}
d^{2} \mathbf{x}_{2}=x_{2} d x_{2} d \phi=J d x_{12} d x_{20} \tag{32}
\end{equation*}
$$

where $\phi$ is the angle between $\mathbf{x}_{20}$ and $\mathbf{x}_{10}$. Inserting an extra factor of 2 to account for the $0<\phi<\pi$ as well as the $\pi<\phi<2 \pi$ domain,


Figure 11: Diagrams for two gluon emission

$$
\begin{equation*}
J\left(x_{12}, x_{02}\right)=\frac{4 x_{21} x_{20}}{\sqrt{\left[\left(x_{21}+x_{20}\right)^{2}-x_{10}^{2}\right]\left[x_{10}^{2}-\left(x_{21}-x_{20}\right)^{2}\right]}} \tag{33}
\end{equation*}
$$

For the 2 gluon emitted squared wavefunction, the second gluon can be emitted from either the $x_{02}$ dipole (lefthand picture in figure 11) or the $x_{12}$ dipole (righthand picture in figure 11). Given these two possibilities, the 2 gluon squared wavefunction can then be written,

$$
\begin{align*}
\Phi^{(2)}\left(\mathbf{x}_{1}, z_{1}\right) & =\left(\frac{\alpha N_{c}}{2 \pi^{2}}\right)^{2} \int d^{2} \mathbf{x}_{2} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} \int d^{2} \mathbf{x}_{3} \int_{z_{0}}^{z_{1}} \frac{d z_{3}}{z_{3}}\left(\frac{x_{02}^{2}}{x_{30}^{2} x_{32}^{2}}+\frac{x_{12}^{2}}{x_{31}^{2} x_{32}^{2}}\right) \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) \\
& =\left(\frac{\alpha N_{c}}{2 \pi^{2}}\right)^{2} \ln ^{2}\left(\frac{z_{1}}{z_{0}}\right) \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) \int d^{2} \mathbf{x}_{2} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} \int d^{2} \mathbf{x}_{3}\left(\frac{x_{02}^{2}}{x_{30}^{2} x_{32}^{2}}+\frac{x_{12}^{2}}{x_{31}^{2} x_{32}^{2}}\right) \tag{34}
\end{align*}
$$

Performing the transform of coordinates (32) using the Jacobian (33), we can also write this solution as

$$
\begin{align*}
= & \left(\frac{2 \alpha N_{c}}{\pi^{2}}\right)^{2} \ln ^{2}\left(\frac{z_{1}}{z_{0}}\right) \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) x_{10}^{2} \int d x_{20} d x_{21} \frac{J\left(x_{20}, x_{21}\right)}{x_{20}^{2} x_{21}^{2}} \\
& \times\left[\int d x_{30} d x_{32} \frac{J\left(x_{30}, x_{32}\right) x_{02}^{2}}{x_{30}^{2} x_{32}^{2}}+\int d x_{32} d x_{31} \frac{J\left(x_{31}, x_{32}\right) x_{12}^{2}}{x_{31}^{2} x_{32}^{2}}\right] \tag{35}
\end{align*}
$$

Now that we have calculated the squared wavefunctions for 1 and 2 soft gluons, we are prepared to generalize to $n$ gluons through the use of a generating functional. Let $\Phi\left(\mathbf{x}_{1}, z_{1}, u(\mathbf{x}, z)\right)$ be defined by the equation,

$$
\begin{align*}
& \left.\frac{\delta}{\delta u\left(\mathbf{x}_{2}, z_{2}\right)} \frac{\delta}{\delta u\left(\mathbf{x}_{3}, z_{3}\right)} \cdots \frac{\delta}{\delta u\left(\mathbf{x}_{n+1}, z_{n+1}\right)} \Phi\left(\mathbf{x}_{1}, z_{1}, u(\mathbf{x}, z)\right)\right|_{u=0} \\
& \quad=\Phi^{(n)}\left(\mathbf{x}_{1}, \mathbf{x}_{2}, \cdots, \mathbf{x}_{n+1} ; z_{2} \cdots z_{n+1}\right) \tag{36}
\end{align*}
$$

where $\Phi^{(n)}$ is the $n$ gluon squared wavefunction, and $\mathbf{x}_{n+1}, z_{n+1}$ are the transverse position and momentum fraction, respectively, of the $n$th gluon. Let us now define the generating functional $Z$ by

$$
\begin{equation*}
\Phi\left(\mathbf{x}_{1}, z_{1}, u\right)=\Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) Z\left(\mathbf{x}_{1}, \mathbf{x}_{0}, z_{1}, u\right) \tag{37}
\end{equation*}
$$

such that the following holds:
$Z\left(\mathbf{x}_{1}, \mathbf{x}_{0}, z_{1}, u\right)=1+\frac{\alpha N_{c}}{2 \pi^{2}} x_{01}^{2} \int d^{2} \mathbf{x}_{2} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} \int_{z_{0}}^{z_{1}} u\left(\mathbf{x}_{2}, z_{2}\right) Z\left(\mathbf{x}_{2}, \mathbf{x}_{1}, z_{2}, u\right) Z\left(\mathbf{x}_{2}, \mathbf{x}_{0}, z_{2}, u\right)$

Using the standard rules for functional differentiation,

$$
\begin{equation*}
\frac{\delta}{\delta u(\mathbf{x})} u(\mathbf{y})=\delta^{(2)}(\mathbf{x}-\mathbf{y}) \quad \frac{\delta}{\delta u(\mathbf{x})} \int d^{2} \mathbf{y} u(\mathbf{y}) f(\mathbf{y})=f(\mathbf{x}) \tag{39}
\end{equation*}
$$

we can demonstrate (36) by reproducing the 2 gluon squared wavefunction (34). Let us calculate the LHS of (36) before setting $u=0$.
$\frac{\delta}{\delta u\left(\mathbf{x}_{3}, z_{3}\right)} \frac{\delta}{\delta u\left(\mathbf{x}_{2}, z_{2}\right)} \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) Z\left(\mathbf{x}_{1}, \mathbf{x}_{0}, z_{1}, u\right)$

$$
\begin{gather*}
=\Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) \frac{\delta}{\delta u\left(\mathbf{x}_{3}, z_{3}\right)} \frac{\delta}{\delta u\left(\mathbf{x}_{2}, z_{2}\right)}\left[1+\frac{\alpha N_{c}}{2 \pi^{2}} \int d^{2} \mathbf{x}_{\alpha} \frac{x_{10}^{2}}{x_{\alpha 0}^{2} x_{\alpha 1}^{2}}\right. \\
\left.\times \int_{z_{0}}^{z_{1}} \frac{d z_{\alpha}}{z_{\alpha}} u\left(\mathbf{x}_{\alpha}, z_{\alpha}\right) Z\left(\mathbf{x}_{\alpha}, \mathbf{x}_{1}, z_{\alpha}, u\right) Z\left(\mathbf{x}_{\alpha}, \mathbf{x}_{0}, z_{\alpha}, u\right)\right]  \tag{40}\\
=\frac{\alpha N_{c}}{2 \pi^{2}} \frac{1}{z_{2}} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right) \frac{\delta}{\delta u\left(\mathbf{x}_{3}, z_{3}\right)}\left[Z\left(\mathbf{x}_{2}, \mathbf{x}_{1}, z_{2}, u\right) Z\left(\mathbf{x}_{2}, \mathbf{x}_{0}, z_{2}, u\right)\right] \\
=\frac{\alpha N_{c}}{2 \pi^{2}} \frac{1}{z_{2}} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right)\left[\frac{\delta Z\left(\mathbf{x}_{2}, \mathbf{x}_{1}, z_{2}, u\right)}{\delta u\left(\mathbf{x}_{3}, z_{3}\right)} Z\left(\mathbf{x}_{2}, \mathbf{x}_{0}, z_{2}, u\right)\right. \\
\left.\quad+\frac{\delta Z\left(\mathbf{x}_{2}, \mathbf{x}_{0}, z_{2}, u\right)}{\delta u\left(\mathbf{x}_{3}, z_{3}\right)} Z\left(\mathbf{x}_{2}, \mathbf{x}_{1}, z_{2}, u\right)\right]  \tag{41}\\
=\left(\frac{\alpha N_{c}}{2 \pi^{2}}\right)^{2} \frac{1}{z_{2}} \frac{1}{z_{3}} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} \Phi^{(0)}\left(\mathbf{x}_{1}, z_{1}\right)\left[\frac{x_{21}^{2}}{x_{23}^{2} x_{31}^{2}} Z_{23 ; 3} Z_{13 ; 3} Z_{20 ; 2}\right.  \tag{42}\\
\left.\quad+\frac{x_{20}^{2}}{x_{23}^{2} x_{30}^{2}} Z_{23 ; 3} Z_{31 ; 3} Z_{21 ; 2}\right]
\end{gather*}
$$

where hopefully the abbreviated notation for $Z_{\alpha \beta ; \gamma}:=Z\left(\mathbf{x}_{\alpha}, \mathbf{x}_{\beta}, z_{\gamma}, u\right)$ is clear. Now letting $u=0$ in (42) so that $Z_{\alpha \beta ; \gamma}=1$, and taking the appropriate integrals, we obtain (34).

While (38) yields the $n$ gluon squared wavefunctions upon functional differentiation, it fails to address virtual corrections and does not satisfy

$$
\begin{equation*}
\left.\int d^{2} \mathbf{x}_{1} \int_{0}^{1} d z_{1} \Phi\left(\mathbf{x}_{1}, z_{1}, u\right)\right|_{u=1}=1 \tag{43}
\end{equation*}
$$

Cutting off the ultraviolet divergences caused by $x_{20}$ or $x_{21}$ going to zero, we introduce a size cutoff $\rho \ll R_{\text {target }}$ such that $x_{20}, x_{21} \geq \rho$. By enforcing (43) at each order in $\alpha$ one can obtain the generating functional with virtual corrections,
$Z\left(\mathrm{x}_{1}, \mathrm{x}_{0}, z_{1}, u\right)$

$$
\begin{align*}
= & \exp \left[-\frac{2 \alpha N_{c}}{\pi} \ln \left(\frac{x_{10}}{\rho}\right) \ln \left(\frac{z_{1}}{z_{0}}\right)\right]+\frac{\alpha N_{c}}{2 \pi^{2}} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \\
& \times \int_{\rho} \exp \left[-\frac{2 \alpha N_{c}}{\pi} \ln \left(\frac{x_{10}}{\rho}\right) \ln \left(\frac{z_{1}}{z_{2}}\right)\right] \frac{d^{2} \mathbf{x}_{2} x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} u\left(\mathbf{x}_{2}, z_{2}\right) Z_{2,1 ; 2} Z_{2,0 ; 2} \tag{44}
\end{align*}
$$

This equation represents a classical branching process and is exact in the leading $\operatorname{logarithmic~approximation.~Another~form~of~this~equation~we~will~use,~letting~} Y:=$ $\ln \left(\frac{z_{1}}{z_{0}}\right), y:=\ln \left(\frac{z_{2}}{z_{0}}\right)$, and $\bar{\alpha}:=\frac{\alpha N_{c}}{\pi}$, is
$Z\left(\mathbf{x}_{1}, \mathbf{x}_{0}, z_{1}, u\right)$

$$
\begin{align*}
= & \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) Y\right]+\frac{\bar{\alpha}}{2 \pi} \int_{z_{0}}^{z_{1}} \frac{d z_{2}}{z_{2}} \\
& \times \int_{\rho} \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)(Y-y)\right] \frac{d^{2} \mathbf{x}_{2} x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} u\left(\mathbf{x}_{2}, z_{2}\right) Z_{2,1 ; 2} Z_{2,0 ; 2} \tag{45}
\end{align*}
$$

### 3.2.3 BFKL from the $n$ gluon onium wavefunction

The generating functional in (45) can now be rewritten as an amplitude. Adding the two equal terms at first order in $\bar{\alpha}$ yields a factor of two in second term of the RHS below:

$$
T\left(x_{10}, z_{1} ; Q, z\right)
$$

$$
\begin{align*}
= & \bar{\alpha} v\left(Q, x_{10}\right) \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) Y\right]+2 \bar{\alpha} \int_{z}^{z_{1}} \frac{d z_{2}}{z_{2}} \\
& \times \int_{\rho} \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)(Y-y)\right] \tilde{K}\left(x_{10}, x_{12}\right) d x_{12} T\left(x_{12}, z_{2} ; Q, z\right) \tag{46}
\end{align*}
$$

where

$$
\begin{equation*}
\tilde{K}\left(x_{10}, x_{12}\right)=\frac{1}{2 \pi} \int_{\rho} \frac{x_{10}^{2}}{x_{20}^{2} x_{21}^{2}} J\left(x_{21}, x_{20}\right) d x_{20} \tag{47}
\end{equation*}
$$

Let us now write $T\left(Y, Q x_{10}\right):=T\left(x_{10}, z_{1} ; Q, z\right)$ as the (inverse) Mellin transform ${ }^{3}$ of $T_{\omega}\left(Q x_{10}\right)$.

$$
\begin{equation*}
T\left(Y, Q x_{10}\right)=\int_{c-i \infty}^{c+i \infty} \frac{d \omega}{2 \pi i} e^{\omega Y} T_{\omega}\left(Q x_{10}\right) \tag{48}
\end{equation*}
$$

This contour integral is a vertical line in the complex plane drawn such that $c$ is greater than the real part of any singularities of $T_{\omega}$. Note that the first term on the RHS of (46) can be written as

$$
\begin{equation*}
\bar{\alpha} v\left(Q, x_{10}\right) \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) Y\right]=\int_{c-i \infty}^{c+i \infty} \frac{d \omega}{2 \pi i} e^{\omega Y} \frac{\bar{\alpha} v\left(Q, x_{10}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)} \tag{49}
\end{equation*}
$$

since the pole of $\omega=-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)$ leads to the residue on the LHS of the equation. Evaluating the second term on the RHS of (46),

$$
\begin{aligned}
& 2 \bar{\alpha} \int_{z}^{z_{1}} \frac{d z_{2}}{z_{2}} \int_{\rho} \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)(Y-y)\right] \tilde{K}\left(x_{10}, x_{12}\right) d x_{12} T\left(x_{12}, z_{2} ; Q, z\right) \\
& =2 \bar{\alpha} \int_{0}^{Y} d y \int_{\rho} \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)(Y-y)\right] \tilde{K}\left(x_{10}, x_{12}\right) d x_{12} \int_{c-i \infty}^{c+i \infty} \frac{d \omega}{2 \pi i} e^{\omega(y-Y)} T_{\omega}\left(Q x_{12}\right) \\
& =2 \bar{\alpha} \int_{c-i \infty}^{c+i \infty} \frac{d \omega}{2 \pi i} \int_{\rho} \tilde{K}\left(x_{10}, x_{12}\right) d x_{12} \frac{1}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)} \\
& \times\left\{1-\exp \left[-\left(2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)+\omega\right) Y\right]\right\} T_{\omega}\left(Q x_{12}\right)
\end{aligned}
$$

[^2]\[

$$
\begin{equation*}
\approx 2 \bar{\alpha} \int_{c-i \infty}^{c+i \infty} \frac{d \omega}{2 \pi i} \int_{\rho} d x_{12} \frac{\tilde{K}\left(x_{10}, x_{12}\right) T_{\omega}\left(Q x_{12}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)} \tag{50}
\end{equation*}
$$

\]

Where we took the leading order of the term in curly braces in the last step. Using (49) and (50), we now see that in Mellin space, (46) takes the following form:

$$
\begin{equation*}
T_{\omega}\left(Q x_{10}\right)=\bar{\alpha} \frac{v\left(Q x_{10}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)}+2 \bar{\alpha} \int d x_{12} \frac{\tilde{K}\left(x_{10}, x_{12}\right) T_{\omega}\left(Q x_{12}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)} \tag{51}
\end{equation*}
$$

Notice that if we redefine the kernel as

$$
\begin{equation*}
K\left(x_{10}, x_{12}\right):=\tilde{K}\left(x_{10}, x_{12}\right)-\delta\left(x_{10}-x_{12}\right) \ln \left(\frac{x_{10}}{\rho}\right) \tag{52}
\end{equation*}
$$

then (51) takes on a particularly simple form.

$$
\begin{gather*}
T_{\omega}\left(Q x_{10}\right)=\bar{\alpha} \frac{v\left(Q x_{10}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)}+2 \bar{\alpha} \int d x_{12}\left[\frac{K\left(x_{10}, x_{12}\right) T_{\omega}\left(Q x_{12}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)}\right.  \tag{53}\\
\left.+\frac{\delta\left(x_{10}-x_{12}\right) \ln \left(\frac{x_{10}}{\rho}\right) T_{\omega}\left(Q x_{12}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)}\right] \\
T_{\omega}\left(Q x_{10}\right)\left(\frac{\omega}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)}\right)=\bar{\alpha} \frac{v\left(Q x_{10}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)}+2 \bar{\alpha} \int d x_{12} \frac{K\left(x_{10}, x_{12}\right) T_{\omega}\left(Q x_{12}\right)}{\omega+2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)}  \tag{54}\\
T_{\omega}\left(Q x_{10}\right)=\frac{\bar{\alpha}}{\omega} v\left(Q x_{10}\right)+\frac{2 \bar{\alpha}}{\omega} \int d x_{12} K\left(x_{10}, x_{12}\right) T_{\omega}\left(x_{12} Q\right) \tag{55}
\end{gather*}
$$

This is, in fact, the celebrated BFKL equation. Let us now show that it yields the well known eigenvalue $\chi(\lambda)=\psi(1)-\frac{1}{2} \psi(1-\lambda / 2)-\frac{1}{2} \psi(\lambda / 2)$, with $\psi(x):=\frac{d}{d x} \ln \Gamma(x)$ being the digamma function and $\psi(1)=\gamma_{e}$ Euler's constant. Let us first manipulate the $\tilde{K}$ part of the kernel in (52). Recalling (32) and (33),

$$
\begin{align*}
\tilde{K}\left(x_{10}, x_{12}\right) & =\frac{1}{2 \pi} \int_{\rho}^{\infty} \frac{x_{10}^{2}}{x_{12}^{2} x_{20}^{2}} J\left(x_{21}, x_{20}\right) d x_{20} \\
& =\frac{2 x_{10}^{2}}{\pi x_{12}} \int_{\rho}^{\infty} \frac{d x_{20}}{x_{20}} \frac{1}{\sqrt{\left[\left(x_{21}+x_{20}\right)^{2}-x_{10}^{2}\right]\left[x_{10}^{2}-\left(x_{21}-x_{20}\right)^{2}\right]}} \tag{56}
\end{align*}
$$

Bringing in an identity that relates transverse lengths and Bessel functions,

$$
\begin{gather*}
\frac{\pi}{2} \int_{0}^{\infty} b d b J_{0}\left(b x_{01}\right) J_{0}\left(b x_{20}\right) J_{0}\left(b x_{12}\right)=\frac{1}{\sqrt{\left[\left(x_{21}+x_{20}\right)^{2}-x_{10}^{2}\right]\left[x_{10}^{2}-\left(x_{21}-x_{20}\right)^{2}\right]}}  \tag{57}\\
\tilde{K}\left(x_{10}, x_{12}\right)=\frac{x_{10}^{2}}{x_{12}} \int_{0}^{\infty} b d b J_{0}\left(b x_{01}\right) J_{0}\left(b x_{12}\right) \int_{\rho}^{\infty} \frac{d x_{20}}{x_{20}} J_{0}\left(b x_{20}\right) \tag{58}
\end{gather*}
$$

Let us tackle the $x_{20}$ integral:

$$
\begin{align*}
\int_{\rho}^{\infty} \frac{d x_{20}}{x_{20}} J_{0}\left(b x_{20}\right) & =\lim _{y \rightarrow 0}\left[\int_{0}^{\infty} d x_{20} x^{y-1} J_{0}\left(b x_{20}\right)-\int_{0}^{\rho} d x_{20} x^{y-1} J_{0}\left(b x_{20}\right)\right]  \tag{59}\\
& =\lim _{y \rightarrow 0} 2^{y-1} b^{-y} \frac{\Gamma\left(\frac{y}{2}\right)}{\Gamma\left(1-\frac{y}{2}\right)}-\frac{\rho^{y}}{y} \\
& =\lim _{y \rightarrow 0}\left(\frac{2}{b}\right)^{y} \frac{\Gamma\left(\frac{y}{2}+1\right)}{y \Gamma\left(1-\frac{y}{2}\right)}-\frac{\rho^{y}}{y} \\
& =\lim _{y \rightarrow 0} \frac{\left(\frac{2}{b}\right)^{y}}{\Gamma\left(1-\frac{y}{2}\right)} \frac{\Gamma\left(\frac{y}{2}+1\right)-\Gamma(1)}{y}+\frac{2^{y} b^{-y}}{\Gamma\left(1-\frac{y}{2}\right)} \frac{\Gamma(1)}{y}-\frac{\rho^{y}}{y} \\
& =\lim _{y \rightarrow 0} \frac{\left(\frac{2}{b}\right)^{y}}{\Gamma\left(1-\frac{y}{2}\right)} \frac{\Gamma\left(\frac{y}{2}+1\right)-\Gamma(1)}{y}+\frac{1}{\Gamma\left(1-\frac{y}{2}\right)} \frac{\left(\frac{2}{b}\right)^{y}-\rho^{y}}{y} \\
& =\psi(1)-\ln \frac{b \rho}{2} \tag{60}
\end{align*}
$$

Note that in the above integral we used the standard formula $\Gamma(x)=\Gamma(x+1) / x$. In (59), we also used the approximation $J_{0}\left(b x_{20}\right) \approx 1$ in the second term (red), as its argument is bounded by $\rho$. The integral in the first term (blue) is given in Gradshteyn and Ryzhik [9], p. 668, 6.516-14.
Using this result, let us now evince the eigenvalue for the eigenfunction $x_{12}^{\lambda}$ of the kernel $K$ in (55). Although Mueller omits this derivation in [11] due to it being
"straightforward", it is still quite a bit of work to show. Given the importance of the BFKL eigenvalue, we will perform the full calculation. To do so, we will make use of the Taylor series for Bessel functions,

$$
\begin{equation*}
J_{0}\left(b x_{12}\right)=\sum_{m=0}^{\infty} \frac{(-1)^{m}}{(m!)^{2}}\left(\frac{b x_{12}}{2}\right)^{2 m} \tag{61}
\end{equation*}
$$

Other techniques used will be summarized below.

$$
\begin{align*}
& \int d x_{12} K\left(x_{10}, x_{12}\right) x_{12}^{\lambda} \\
& =\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda}\left[\frac{x_{10}^{2}}{x_{12}} \int_{0}^{\infty} b d b J_{0}\left(b x_{01}\right) J_{0}\left(b x_{12}\right)\left(\psi(1)-\ln \frac{b \rho}{2}\right)-\delta\left(x_{10}-x_{12}\right) \ln \left(\frac{x_{10}}{\rho}\right)\right]  \tag{62}\\
& =\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}}\left\{\int_{0}^{\infty} b d b J_{0}\left(b x_{01}\right) J_{0}\left(b x_{12}\right)\left(\psi(1)-\ln \frac{b x_{10}}{2}\right)\right\} \\
& =\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}} \sum_{m=0}^{\infty} \frac{(-1)^{m}}{(m!)^{2}} \int_{0}^{\infty} b d b\left(\psi(1)-\ln \frac{b x_{10}}{2}\right)\left(\frac{b x_{01}}{2}\right)^{2 m} J_{0}\left(b x_{12}\right) \\
& =\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}} \sum_{m=0}^{\infty} \frac{(-1)^{m}}{(m!)^{2}}\left(\frac{m!}{x_{12}^{2} \Gamma(-m)} \psi(1)-\frac{\partial}{\partial(2 m)}\right)\left[\left(\frac{x_{10}}{2}\right)^{2 m} \int_{0}^{\infty} b d b J_{0}\left(b x_{12}\right) b^{2 m}\right] \tag{63}
\end{align*}
$$

$=\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}^{3}} \sum_{m=0}^{\infty} \frac{(-1)^{m}}{(m!)^{2}}\left(\frac{m!}{\Gamma(-m)} \psi(1)-\frac{\partial}{\partial(2 m)}\right)\left[2\left(\frac{x_{10}}{x_{12}}\right)^{2 m} \frac{\Gamma(m+1)}{\Gamma(-m)}\right]$

$$
\begin{equation*}
=\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}^{3}} \sum_{m=0}^{\infty} \frac{(-1)^{m}}{(m!)^{2}} 2\left(\frac{x_{10}}{x_{12}}\right)^{2 m}\left(\frac{m!}{\Gamma(-m)} \psi(1)-\ln \left(\frac{x_{10}}{x_{12}}\right) \frac{\Gamma(m+1}{\Gamma(-m)}\right. \tag{64}
\end{equation*}
$$

$$
\left.-\frac{\partial}{2 \partial m} \frac{\Gamma(m+1)}{\Gamma(-m)}\right)
$$

$$
\begin{align*}
= & \int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}^{3}} \sum_{m=0}^{\infty} \frac{(-1)^{m}}{(m!)^{2}} 2\left(\frac{x_{10}}{x_{12}}\right)^{2 m}\left(\frac{m!}{\Gamma(-m)} \psi(1)\right. \\
& \left.-\frac{1}{2} \frac{\Gamma^{\prime}(m+1) \Gamma(-m)+\Gamma^{\prime}(-m) \Gamma(m+1)}{\Gamma^{2}(-m)}\right) \\
= & \int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}^{3}} \sum_{m=0}^{\infty} \frac{(-1)^{m}}{(m!)^{2}} \frac{m!}{\Gamma(-m)}\left(\frac{x_{10}}{x_{12}}\right)^{2 m} 2\left[\psi(1)-\frac{1}{2} \psi(m+1)-\frac{1}{2} \psi(-m)\right]  \tag{65}\\
& =\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}^{3}} \sum_{m=0}^{\infty} \frac{(-1)^{m}}{m!} \frac{1}{\Gamma(-m)}\left(\frac{x_{10}}{x_{12}}\right)^{2 m} 2 \chi(-2 m) \tag{66}
\end{align*}
$$

$$
=\int_{\rho}^{\infty} d x_{12} x_{12}^{\lambda} \frac{x_{10}^{2}}{x_{12}^{3}} x_{01} \delta\left(x_{01}-x_{12}\right) \chi\left(\partial / \partial x_{12}\right)
$$

$$
\begin{equation*}
\int d x_{12} K\left(x_{10}, x_{12}\right) x_{12}^{\lambda}=\chi(\lambda) x_{01}^{\lambda} \tag{67}
\end{equation*}
$$

In (62) (green) we used the orthogonality/closure relation for Bessel functions:

$$
\begin{equation*}
\int_{0}^{\infty} b d b J_{0}\left(b x_{01}\right) J_{0}\left(b x_{12}\right)=\frac{1}{x_{12}} \delta\left(x_{10}-x_{12}\right) \tag{68}
\end{equation*}
$$

In (63) (blue) we used the same Gradshteyn and Ryzhik integral as in (59). In 64) (red) and 66) (cyan) we used

$$
\begin{equation*}
\sum_{m=0}^{\infty} \frac{(-1)^{m}}{m!} \frac{1}{\Gamma(-m)}\left(\frac{x_{10}}{x_{12}}\right)^{2 m}=\frac{x_{10}}{2} \delta\left(x_{10}-x_{12}\right) \tag{69}
\end{equation*}
$$

causing the red term to drop out. In (63) (red) we used (68) and 69). Finally, in (65) (magenta) we defined

$$
\begin{equation*}
\chi(\lambda):=\psi(1)-\frac{1}{2} \psi\left(1-\frac{\lambda}{2}\right)-\frac{1}{2} \psi\left(\frac{\lambda}{2}\right) \tag{70}
\end{equation*}
$$

### 3.3 The Pomeron from BFKL

Now in possession of the BFKL eigenvalue equation (67), we may demonstrate the emergence of the Pomeron. Let us begin by inverse Mellin transforming the amplitude once again.

$$
\begin{equation*}
T_{\omega}\left(x_{12}, Q\right)=\int_{c-i \infty}^{c+i \infty} \frac{d \lambda}{2 \pi i}\left(Q x_{12}\right)^{\lambda} T_{\lambda \omega} \tag{71}
\end{equation*}
$$

The BFKL equation (55) can be easily solved for $T_{\lambda \omega}$.

$$
\begin{gather*}
\int_{c-i \infty}^{c+i \infty} \frac{d \lambda}{2 \pi i}\left[T_{\omega}(\omega-2 \bar{\alpha} \chi(\lambda))\right]=\int_{c-i \infty}^{c+i \infty} \frac{d \lambda}{2 \pi i} \bar{\alpha} v_{\lambda}\left(Q x_{10}\right)  \tag{72}\\
T_{\lambda \omega}=\frac{\bar{\alpha} v_{\lambda}}{\omega-2 \bar{\alpha} \chi(\lambda)} \tag{73}
\end{gather*}
$$

where $v_{\lambda}$ is the Mellin transform of $v\left(Q x_{10}\right)$. Recalling (48), let us perform the inverse Laplace and inverse Mellin transforms on (73) to solve for the amplitude as a function of energy, where we expect pomeron behavior to manifest itself.

$$
\begin{equation*}
T\left(Y, Q x_{10}\right)=\int_{c-i \infty}^{c+i \infty} \frac{d \omega}{2 \pi i} e^{\omega Y} \int_{c-i \infty}^{c+i \infty} \frac{d \lambda}{2 \pi i}\left(Q x_{10}\right)^{\lambda} \frac{\bar{\alpha} v_{\lambda}}{\omega-2 \bar{\alpha} \chi(\lambda)} \tag{74}
\end{equation*}
$$

The $\omega$ integral is a simple residue.

$$
\begin{equation*}
T\left(Y, Q x_{10}\right)=\bar{\alpha} \int_{c-i \infty}^{c+i \infty} \frac{d \lambda}{2 \pi i} v_{\lambda} e^{2 \bar{\alpha} \chi(\lambda) Y+\lambda \ln \left(Q x_{10}\right)} \tag{75}
\end{equation*}
$$

Assuming that a) $\ln \left(Q x_{10}\right) \ll \bar{\alpha} Y$, or that the transverse momentum is not too large, and b) $v_{\lambda}$ is a slowly varying function, the integral in (75) can be approximated by the saddle point method. This method evaluates the integral where the phase is approximately stationary. We can see where this occurs by examining the graph of $\chi(\lambda)$ shown on figure 12 .
Let us use the expansion of $\chi(\lambda)$ around $\lambda=1$ [40]:


Figure 12: Graph of $\chi(\lambda)$ between $0<\lambda<2$. Note the saddle point at $\lambda=1$.

$$
\begin{align*}
\chi(\lambda) & \approx 2 \ln 2+\frac{7}{4} \zeta(3)(\lambda-1)^{2}  \tag{76}\\
\chi^{\prime}(\lambda) & \approx \frac{7}{2} \zeta(3)(\lambda-1)  \tag{77}\\
\chi^{\prime \prime}(\lambda) & \approx \frac{7}{2} \zeta(3) \tag{78}
\end{align*}
$$

where $\zeta(x)$ is the Riemann zeta function. The saddle point approximation can be written as

$$
\begin{equation*}
\int_{c-i \infty}^{c+i \infty} d \lambda e^{f(\lambda)-\lambda \bar{x}} \approx \frac{1}{\sqrt{2 \pi f^{\prime \prime}\left(\lambda_{s}\right)}} \exp \left(f\left(\lambda_{s}\right)-\lambda_{s} \bar{x}-\frac{\left[f^{\prime}\left(\lambda_{s}\right)-\bar{x}\right]^{2}}{2 f^{\prime \prime}\left(\lambda_{s}\right)}\right) \tag{79}
\end{equation*}
$$

Applying this approximation to (75), along with (76), (77), and (78), we obtain

$$
\begin{align*}
T\left(Y, Q x_{10}\right) \approx & \frac{\bar{\alpha} v_{1}}{\sqrt{14 \bar{\alpha} \pi \zeta(3) Y}} \exp \left(4 \bar{\alpha} \ln (2) Y-\ln \left(Q x_{10}\right)-\frac{\ln ^{2}\left(Q x_{10}\right)}{14 \bar{\alpha} \zeta(3) Y}\right) \\
& =\frac{\bar{\alpha} v_{1}\left(Q x_{10}\right)}{\sqrt{14 \bar{\alpha} \pi \zeta(3) Y}} e^{\left(\alpha_{\mathbb{P}}-1\right) Y} \exp \left(-\frac{\ln ^{2}\left(Q x_{10}\right)}{14 \bar{\alpha} \zeta(3) Y}\right) \tag{80}
\end{align*}
$$

with

$$
\begin{equation*}
\alpha_{\mathbb{P}}-1=4 \bar{\alpha} \ln (2) \tag{81}
\end{equation*}
$$

By (22), we see that BFKL evolution in the dipole picture indeed leads to the same hard pomeron behavior as in (16).

## 4 The BK Equation and Traveling Wave Solutions

### 4.1 Unitarity corrections to the BFKL equation; the BK equation

What are the consequences of a cross-sectional rise that goes like $e^{\alpha_{\mathbb{P}}-1}$ using (81)? Let us do a quick calculation: let $Q^{2} \approx 10 \mathrm{GeV}$, a moderate value that does not violate the condition under (75). Using the well known formula of Gross, Politzer, and Wilczek for asymptotic freedom [53, 54],

$$
\begin{equation*}
\alpha_{s}(Q)=\frac{2 \pi}{b_{0} \ln (Q / \Lambda)}, \quad b_{0}=11-\frac{2}{3} n_{f} \tag{82}
\end{equation*}
$$

Using $n_{f}=3$ light quarks and $\Lambda=.2 \mathrm{GeV}$, we obtain $\alpha_{s}=.178$. Then, with $N_{c}=3$,

$$
\alpha_{\mathbb{P}}-1=4 \bar{\alpha} \ln (2)=\frac{12 \alpha_{s}}{\pi} \ln (2) \approx .47
$$

Unlike for Reggeons (mesons $\rho, \omega, f_{2}, a_{2}$,etc.) with a Regge trajectory intercept of ${ }^{1}$ $\alpha(0)-1 \approx-.45$, the BFKL pomeron, also called the hard pomeron, causes the crosssection to rise with $s$. This is actually necessary to fit available data, but with such a large power the Froissart-Martin bound [55] (a consequence of unitarity),

$$
\begin{equation*}
\sigma_{t o t}(s)<\frac{\pi}{m_{\pi}^{2}} \ln ^{2}\left(\frac{s}{s_{0}}\right) \tag{83}
\end{equation*}
$$

is violated even within HERA's energy range. It is possible to introduce next to leading order (NLO) corrections to the BFKL equation that allow HERA data to be successfully fit [1], but even these are not enough to tame the eventual rise predicted by the LO BFKL equation ${ }^{5}$. A great deal of effort throughout the 90 s went into formulating QCD evolution equations that preserve unitarity. This led to the BJIMWLK equations [41, 56, 57], which were several different techniques: a functional renormalization group equation, an infinite hierarchy of coupled integro-differential equations, and a Langevin equation. In 1999, Kovchegov managed to considerably

[^3]simplify Balitsky's equation using Mueller's dipole formulation, deriving what is now known as the BK equation. We will review the presentation of [39, 40] condensing and simplifying notation where possible.
Following [12] [13, we will implement a dipole number density $n\left(x_{01}, Y,|\mathbf{b}|, x_{1}\right)$, which when convoluted with the photon dissociation wavefunction squared, $\Phi\left(z_{1}, x_{01}\right)$, gives
\[

$$
\begin{equation*}
N\left(x_{1}, Y\right)=\int d^{2} x_{01} \int_{0}^{1} d z_{1} \Phi\left(z_{1}, x_{01}\right) n\left(x_{01}, Y, x_{1}\right) \tag{84}
\end{equation*}
$$

\]

where $N\left(x_{1}, Y\right)$ is the propagator of the virtual photon through a target nucleus. ${ }^{6}$. The BK equation is usually derived in the frame of the target with the evolution put into probe. We will see (84) obtains when we define $n\left(x_{01}, Y, x_{1}\right)$ by

$$
\begin{equation*}
\frac{1}{2 \pi x_{1}^{2}} n_{1}\left(x_{01}, Y,|\mathbf{b}|, x_{1}\right):=\left.\frac{\delta}{\delta u\left(\mathbf{x}_{1}\right)} Z\left(\mathbf{x}_{01}, Y, u\right)\right|_{u=1} \tag{85}
\end{equation*}
$$

Likewise, we can define the dipole pair density

$$
\begin{equation*}
\frac{1}{2 \pi x_{1}^{2}} \frac{1}{2 \pi x_{2}^{2}} n_{2}\left(x_{01}, Y, x_{1}, x_{2}\right)=\left.\frac{1}{2} \frac{\delta}{\delta u\left(\mathbf{x}_{1}\right)} \frac{\delta}{\delta u\left(\mathbf{x}_{2}\right)} Z\left(\mathbf{x}_{01}, Y, u\right)\right|_{u=1} \tag{86}
\end{equation*}
$$

and generalizing to the group of $k$ dipoles with sizes $x_{1}, \ldots, x_{k}$,

$$
\begin{equation*}
\prod_{i=1}^{k} \frac{1}{2 \pi x_{i}^{2}} n_{k}\left(x_{01}, Y, x_{1}, \ldots, x_{k}\right)=\left.\frac{1}{k!} \prod_{i=1}^{k} \frac{\delta}{\delta u\left(\mathbf{x}_{i}\right)} Z\left(\mathbf{x}_{01}, Y, u\right)\right|_{u=1} \tag{87}
\end{equation*}
$$

The result of multiple functional differentiation in (87) is [39]

$$
\begin{aligned}
& n_{i}\left(x_{01}, Y, \mathbf{x}_{1}, \ldots, \mathbf{x}_{k}\right) \\
& \quad=\frac{\bar{\alpha}}{2 \pi} \int_{0}^{Y} d y \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{01}}{\rho}\right)(Y-y)\right] \int_{\rho} d^{2} \mathbf{x}_{2}^{\prime} \frac{x_{01}^{2}}{x_{02}^{2} x_{12}^{2}} \\
& \left.\quad \times\left[2 n_{i}\left(x_{02}, Y, \mathbf{x}_{1}, \ldots, \mathbf{x}_{k}\right)+\sum_{j+k=i} n_{j}\left(x_{02}, Y, \mathbf{x}_{1}, \ldots, \mathbf{x}_{k}\right) n_{k}\left(x_{12}, Y, \mathbf{x}_{1}, \ldots, \mathbf{x}_{k}\right)\right] 88\right)
\end{aligned}
$$

[^4]The total interaction cross-section is the sum of the interactions of each of the groups of $k$ dipoles with the target. We can write this as

$$
\begin{align*}
N\left(\mathbf{x}_{01}, Y\right)= & \int \frac{d^{2} \mathbf{x}_{1}}{2 \pi x_{1}^{2}} n_{1}\left(x_{01}, Y, \mathbf{x}_{1}\right) \\
& +\int \frac{d^{2} \mathbf{x}_{1}}{2 \pi x_{1}^{2}} \frac{d^{2} \mathbf{x}_{2}}{2 \pi x_{2}^{2}} n_{2}\left(x_{01}, Y, \mathbf{x}_{1}, \mathbf{x}_{2}\right)+\ldots \\
= & \sum_{i=1}^{\infty} \int \frac{d^{2} \mathbf{x}_{1}}{2 \pi x_{1}^{2}} \cdots \frac{d^{2} \mathbf{x}_{i}}{2 \pi x_{2}^{2}} n_{i}\left(x_{01}, Y, \mathbf{x}_{1}, \ldots, \mathbf{x}_{i}\right) \tag{89}
\end{align*}
$$

Performing these operations on (88) yields

$$
\begin{align*}
N\left(\mathbf{x}_{01}, Y\right)= & \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) Y\right]+\frac{\bar{\alpha}}{2 \pi} \int_{0}^{Y} d y \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right)(Y-y)\right] \\
& \times \int_{\rho} d^{2} \mathbf{x}_{2} \frac{x_{01}^{2}}{x_{02}^{2} x_{12}^{2}}\left[2 N\left(\mathbf{x}_{02}, y\right)-N\left(\mathbf{x}_{02}, y\right) N\left(\mathbf{x}_{12}, y\right)\right] \tag{90}
\end{align*}
$$

Finally, taking the derivative of 90 with respect to Y,

$$
\begin{align*}
\frac{\partial N\left(\mathbf{x}_{01}, Y\right)}{\partial Y}= & -2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) Y\right] \\
& \frac{\bar{\alpha}}{2 \pi} \int_{\rho} d^{2} \mathbf{x}_{2} \frac{x_{01}^{2}}{x_{02}^{2} x_{12}^{2}}\left[2 N\left(\mathbf{x}_{02}, Y\right)-N\left(\mathbf{x}_{02}, Y\right) N\left(\mathbf{x}_{12}, Y\right)\right] \tag{91}
\end{align*}
$$

Rewriting the the first term on the RHS to first order in $\bar{\alpha}$ as

$$
\begin{equation*}
-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) \exp \left[-2 \bar{\alpha} \ln \left(\frac{x_{10}}{\rho}\right) Y\right]=-\frac{\bar{\alpha}}{2 \pi} \ln \left(\frac{x_{10}}{\rho}\right) \int_{\rho} d^{2} \mathbf{x}_{2} 4 \pi \delta^{2}\left(\mathbf{x}_{01}-\mathbf{x}_{02}\right) N\left(\mathbf{x}_{02}, Y\right) \tag{92}
\end{equation*}
$$

we can put (91) into a somewhat simpler form.

$$
\begin{equation*}
\frac{\partial N\left(\mathbf{x}_{01}, Y\right)}{\partial Y}=\frac{\bar{\alpha}}{2 \pi} \int_{\rho} d^{2} \mathbf{x}_{2}\left\{\frac{x_{01}^{2}}{x_{02}^{2} x_{12}^{2}}\left[2 N\left(\mathbf{x}_{02}, Y\right)-N\left(\mathbf{x}_{02}, Y\right) N\left(\mathbf{x}_{12}, Y\right)\right]\right. \tag{93}
\end{equation*}
$$

$$
\left.-4 \pi \delta^{2}\left(\mathbf{x}_{01}-\mathbf{x}_{02}\right) \ln \left(\frac{x_{01}}{\rho}\right) N\left(\mathbf{x}_{02}, Y\right)\right\}
$$

Notice that

$$
\begin{align*}
& \int_{\rho} d^{2} \mathbf{x}_{2} \frac{x_{01}^{2}}{x_{02}^{2} x_{12}^{2}}=2(2 \pi) \int_{\rho} d x_{12} x_{12}\left(\frac{x_{01}^{2}}{x_{02}^{2}}\right) \frac{1}{x_{12}^{2}}=4 \pi \int_{\rho} \frac{d x_{12}}{x_{12}}=4 \pi \ln \left(\frac{x_{01}}{\rho}\right)  \tag{94}\\
& \quad=\int_{\rho} d^{2} \mathbf{x}_{2} 4 \pi \delta^{2}\left(\mathbf{x}_{01}-\mathbf{x}_{02}\right) \ln \left(\frac{x_{01}}{\rho}\right)
\end{align*}
$$

where the factor of 2 after the first equality is due to evaluation at the collinear limit near both $\mathbf{x}_{0}$ and $\mathbf{x}_{1}$. If we take $x_{01} \approx x_{02}$, for instance, the second equality of (94) follows. Using (94), we can write the BK equation in another commonly used form (see [1][2]):

$$
\begin{equation*}
\frac{\partial N\left(\mathbf{x}_{01}, Y\right)}{\partial Y}=\frac{\bar{\alpha}}{2 \pi} \int_{\rho} d^{2} \mathbf{x}_{2} \frac{x_{01}^{2}}{x_{02}^{2} x_{12}^{2}} 2 N\left(\mathbf{x}_{02}, Y\right)-N\left(\mathbf{x}_{01}, Y\right)-N\left(\mathbf{x}_{02}, Y\right) N\left(\mathbf{x}_{12}, Y\right) \tag{95}
\end{equation*}
$$

Aside from the nonlinear product $N\left(\mathbf{x}_{02}, Y\right) N\left(\mathbf{x}_{12}, Y\right)$, this equation is actually the same as the BFKL equation. We can crudely approximate when the solutions to the two equations diverge. Using the fact that the elementary dipole-dipole scattering amplitude is $T^{e l} \sim \alpha^{2}$, the probability of two simultaneous scatterings is $\sim \alpha^{4}$, which is suppressed until the density of dipoles is $n \sim 1 / \alpha^{2}$ (see 128129 for details). At these densities, corrections provided by the nonlinear term are needed to stem the rise of the amplitude. Although the interpretation of this reduction in growth is not completely clear at present-be it due to gluon recombination, color swings, etc.-it must exist to preserve unitarity at high energies. In the t-channel picture, one can view the correction as replacing the single gluon ladder diagram with a "fan diagram" containing triple pomeron vertices, as in figure 13 .

### 4.2 FKPP equation and reaction-diffusion dynamics

In this subsection we will show how the BK equation (95) encodes a branching diffusion of dipoles in the variable $\ln \left(1 / r^{2}\right)$. The equation describing such diffusion is


Figure 13: A fan diagram representing the BK equation in the t-channel.
called the Fisher-Kolmogorov-Petrovsky-Piscounoff (FKPP) equation, which is well known in statistical physics and is equivalent to the BK equation in the aptly named diffusion approximation. We will see the FKPP equation admits a traveling wave solution as dipoles diffuse to smaller sizes with increasing rapidity. The application of the FKPP equation to QCD evolution was first pointed out by Munier and Peschanski in a series of papers in 2003-4 [14, 15, [16].

Starting by Fourier transforming the BK equation (93) and using steps very similar to 62.67), we can rewrite the BK equation for momentum space $\tilde{N}(k, Y)$ using the BFKL eigenvalue $\chi_{\text {Mueller }}(\lambda)^{7}$ we found in (70) as [40]

$$
\begin{equation*}
\frac{\partial \tilde{N}(k, Y)}{\partial Y}=\bar{\alpha} \chi\left(-\frac{\partial}{\partial \ln k^{2}}\right) \tilde{N}(k, Y)-\bar{\alpha} \tilde{N}^{2}(k, Y) \tag{96}
\end{equation*}
$$

Defining $L:=\ln \left(k^{2} / \Lambda_{Q C D}^{2}\right)$,

$$
\begin{equation*}
\frac{\partial \tilde{N}(k, Y)}{\partial Y}=\bar{\alpha} \chi\left(-\partial_{L}\right) \tilde{N}(k, Y)-\bar{\alpha} \tilde{N}^{2}(k, Y) \tag{97}
\end{equation*}
$$

Using a series expansion of $\chi\left(-\partial_{L}\right)$ in the principle branch of the eigenvalue around a point $0<\gamma_{0}<1$,

[^5]\[

$$
\begin{equation*}
\chi\left(-\partial_{L}\right)=\chi\left(\gamma_{0}\right) \mathbf{1}+\chi^{\prime}\left(\gamma_{0}\right)\left(-\partial_{L}-\gamma_{0} \mathbf{1}\right)+\frac{1}{2} \chi^{\prime \prime}\left(\gamma_{0}\right)\left(-\partial_{L}-\gamma_{0} \mathbf{1}\right)^{2}+\frac{1}{6} \chi^{(3)}\left(\gamma_{0}\right)\left(-\partial_{L}-\gamma_{0} \mathbf{1}\right)^{3}+\ldots \tag{98}
\end{equation*}
$$

\]

The diffusion approximation is tantamount to keeping only up to second order terms in (98). Let us work with this truncated series and expand around $\gamma_{0}=\frac{1}{2}$, as we did in the saddle point method used in 3.3.

$$
\begin{equation*}
\chi\left(-\partial_{L}\right) \approx \bar{\chi}\left(-\partial_{L}\right):=\chi\left(\frac{1}{2}\right)+\frac{\chi^{\prime \prime}\left(\frac{1}{2}\right)}{2}\left(\partial_{L}+\frac{1}{2}\right)^{2} \tag{99}
\end{equation*}
$$

If we make the following change of coordinates with $\omega:=\chi\left(\frac{1}{2}\right), D:=\chi^{\prime \prime}\left(\frac{1}{2}\right)$, and $\bar{\gamma}:=1-\frac{1}{2} \sqrt{1+8 \omega / D}$,

$$
\begin{gather*}
t:=\frac{\bar{\alpha} D}{2}(1-\bar{\gamma})^{2} Y  \tag{100}\\
x:=(1-\bar{\gamma})\left(L+\frac{\bar{\alpha} D}{2} Y\right)  \tag{101}\\
u(t, x):=\frac{2}{D(1-\bar{\gamma})^{2}} N\left(\frac{2 t}{\bar{\alpha} D(1-\bar{\gamma})^{2}}, \frac{x}{1-\bar{\gamma}}-\frac{t}{(1-\bar{\gamma})^{2}}\right) \tag{102}
\end{gather*}
$$

then (97) with (99) becomes the FKPP equation:

$$
\begin{equation*}
\partial_{t} u(t, x)=\partial_{x}^{2} u(t, x)+u(t, x)-u^{2}(t, x) \tag{103}
\end{equation*}
$$

This equation is very well studied-see, for example, [45, 46] for comprehensive discussions. To quote from one of those references,

The general goal of our discussion of front propagation into unstable states is to investigate the following front propagation problem: If initially a spatially extended system is in an unstable state everywhere except in some spatially localized region, what will be the large-time dynamical properties and speed of the nonlinear front which will propagate into the unstable state? Are there classes of initial conditions for which the front dynamics converges to some unique asymptotic front state? If so, what characterizes these initial conditions, and what can we say about the asymptotic front properties and the convergence to them? [45]

| Reaction-diffusion | QCD |
| :--- | :--- |
| Occupation fraction $\mathrm{u}(\mathrm{t}, \mathrm{x})$ | Scattering amplitude for the probe off <br>  <br>  <br>  <br>  <br> a frozen realization of the target <br> $T(k, Y)$, or $N(k, Y)$ |
| Average occupation fraction $\langle u(t, x)\rangle$ | Physical scattering amplitude $A=\langle T\rangle$ |
| Space variable $x$, sometimes $L$ | $\ln \left(k^{2} / \Lambda^{2}\right)$ or $\ln \left(1 / r^{2} \Lambda^{2}\right)$ |
| Time variable t | Rapidity $\bar{\alpha} Y$ |
| Average maximum density of particles | $1 / \alpha^{2}$ |
| $N$ |  |
| Position of the front $X(t)$ | Saturation scale $\ln \left(Q_{s}^{2}(Y) / \Lambda^{2}\right)$ |
| Branching-diffusion kernel $\omega\left(-\partial_{x}\right)$, <br> $\left(\omega\left(-\partial_{x}\right)=\partial_{x}^{2}+1\right.$ for FKPP) | BFKL kernel $\chi\left(-\partial_{\ln } k^{2}\right)$ or its <br> equivalent in coordinate space |

Table 1: A dictionary between reaction-diffusion and QCD variables. [1]

Let us turn our attention towards some of these issues. In short, an initial condition $u(0, x)$ will evolve into a traveling wave solution $u(t, x)=u(x-v t)$ with an asymptotic front velocity. Using the known result from FKPP analysis, [15]

$$
\begin{equation*}
u(t, x) \quad t \xrightarrow{\sim} \infty \quad w\left(x-2 t+\frac{3}{2} \ln t\right) \tag{104}
\end{equation*}
$$

and assuming an exponential solution,

$$
\begin{equation*}
u(t, x) \sim \exp \left(x-2 t+\frac{3}{2} \ln t\right) \tag{105}
\end{equation*}
$$

we may use the mappings $(100),(101)$, and $(102)$ to write

$$
\begin{gather*}
N(Y, k) \sim u(t, x) \sim \exp \left\{(1-\bar{\gamma})\left(L+\frac{\bar{\alpha} D}{2} Y\right)-2 \frac{\bar{\alpha} D}{2}(1-\bar{\gamma})^{2} Y+\frac{3}{2} \ln \left[\frac{\bar{\alpha} D}{2}(1-\bar{\gamma})^{2} Y\right]\right\}  \tag{106}\\
=\exp (1-\bar{\gamma}) \exp \left[L+\frac{\bar{\alpha} D}{2} Y-\bar{\alpha} D(1-\bar{\gamma}) Y\right] Y^{\frac{3}{2(1-\bar{\gamma})}}\left(\frac{\bar{\alpha} D}{2}(1-\bar{\gamma})^{2}\right)^{\frac{3}{2(1-\bar{\gamma})}} \\
=k_{0}^{-2} k^{2} \exp \left[-\bar{\alpha} D\left(\frac{1}{2}-\bar{\gamma}\right) Y\right] Y^{\frac{3}{2(1-\bar{\gamma})}}
\end{gather*}
$$



Figure 14: Geometric scaling data: the total cross section $\sigma_{t o t}^{\gamma^{*} p \rightarrow X}$ as a function of $\tau:=Q^{2} / Q_{s}^{2}(x)$ for $x<.01$. 38]

$$
\begin{equation*}
=\frac{k^{2}}{Q_{s}^{2}(Y)}, \quad Q_{s}^{2}(Y)=k_{0}^{2} Y^{-\frac{3}{2(1-\bar{\gamma})}} e^{\bar{\alpha} D\left(\frac{1}{2}-\bar{\gamma}\right) Y} \tag{107}
\end{equation*}
$$

where $k_{0}^{-2}$ absorbs the constants. The result of these manipulations is to demonstrate that

$$
\begin{equation*}
N(Y, k)=N\left(\frac{k^{2}}{Q_{s}^{2}(Y)}\right) \tag{108}
\end{equation*}
$$

which is the definition of geometric scaling, a feature strikingly revealed in the data, as shown in figure 14. Geometric scaling was known before Munier and Peschanski showed it was a consequence of the FKPP (see [35, 36]), but these authors framed the BK equation in the larger context of the universality class of the FKPP equation. In fact, the full BK equation (not using the diffusion approximation) and the NLO BFKL equation have both been shown to be a part of this universality class [1], meaning that all of these equations, details aside, exhibit branching diffusion with a saturation mechanism. This has been one of the pivotal discoveries in QCD over the last decade.

It is possible to analytically determine the velocity of the traveling wave predicted
by the FKPP equation. Because the wavefront mediates between the high density and low density regions in $x$, matching amplitudes at the two conditions allows us to determine a critical condition at the wavefront. This critical condition, in a certain interpretation, then yields the wavefront velocity.

First let us investigate the critical condition using a method explained in [47]. Starting from the BK equation (97), and using the Laplace transform,

$$
\begin{equation*}
N(k, \omega)=\int d Y e^{-\omega Y} N(k, Y) \tag{109}
\end{equation*}
$$

with a proposed ansatz 42]

$$
\begin{equation*}
N(k, \omega)=N(\omega) e^{[\gamma(\omega)-1] L} \tag{110}
\end{equation*}
$$

where $L:=\ln \left(k^{2} / \Lambda^{2}\right)$ as before and $\gamma(\omega)$ is the Mellin space argument of the BFKL eigenvalue (also called the anomalous dimension), we obtain

$$
\begin{equation*}
\omega e^{\omega Y} N(k, \omega)=\bar{\alpha} \chi(\gamma(\omega)) e^{\omega Y} N(k, \omega)-\bar{\alpha} \int_{c-i \infty}^{c+i \infty} \frac{d \omega^{\prime}}{2 \pi i} e^{\left(\omega+\omega^{\prime}\right) Y} N(k, \omega) N\left(k, \omega^{\prime}\right) \tag{111}
\end{equation*}
$$

Shifting $\omega \rightarrow \omega-\omega^{\prime}$ in the integral on the RHS,

$$
\begin{equation*}
[\omega-\bar{\alpha} \chi(\gamma(\omega))] N(\omega) e^{[\gamma(\omega)-1] L}=-\bar{\alpha} \int_{c-i \infty}^{c+i \infty} \frac{d \omega^{\prime}}{2 \pi i} N\left(\omega-\omega^{\prime}\right) N\left(\omega^{\prime}\right) e^{\left[\gamma\left(\omega-\omega^{\prime}\right)+\gamma\left(\omega^{\prime}\right)-2\right] L} \tag{112}
\end{equation*}
$$

We may again use the saddle approximation (79) on the integral on the RHS, approximating around the choice $\omega^{\prime}=\omega / 2$ at which the derivative of the exponent vanishes. We obtain

$$
\begin{equation*}
[\omega-\bar{\alpha} \chi(\gamma(\omega))] N(\omega) e^{[\gamma(\omega)-1] L}=-\frac{\bar{\alpha}}{\sqrt{4 \pi \gamma^{\prime \prime}(\omega / 2) L}} N^{2}\left(\frac{\omega}{2}\right) e^{[2 \gamma(\omega / 2)-2] L} \tag{113}
\end{equation*}
$$

In the region where the density is dilute, the nonlinear RHS is approximately zero, yielding

$$
\begin{equation*}
\text { Dilute Region : } \quad \omega-\bar{\alpha} \chi(\gamma(\omega))=0 \tag{114}
\end{equation*}
$$

On the other hand, we may match exponents in in the saturation region to obtain a different condition.

$$
\begin{equation*}
\text { Saturation Region: } \quad \gamma(\omega)=2 \gamma\left(\frac{\omega}{2}\right)-1 \tag{115}
\end{equation*}
$$

which is satisfied by

$$
\begin{equation*}
\gamma(\omega)=C \omega+1 \tag{116}
\end{equation*}
$$

for some constant $C$. We may solve for $C$ using the derivative of (116) to obtain

$$
\begin{align*}
\gamma(\omega) & =\gamma^{\prime}(\omega) \omega+1 \\
\gamma^{\prime}(\omega) & =\frac{\gamma(\omega)-1}{\omega} \tag{117}
\end{align*}
$$

Taking the derivative of the dilute condition (114),

$$
\begin{equation*}
\bar{\alpha} \chi^{\prime}(\gamma)=\frac{1}{\gamma^{\prime}(\omega)} \tag{118}
\end{equation*}
$$

Finally, we expect (118) to match with (117) at some critical value $\gamma_{c}=\gamma\left(\omega_{c}\right)$ at the wavefront where the dilute and saturation regions meet. Thus we obtain

$$
\begin{align*}
\bar{\alpha} \chi^{\prime}\left(\gamma_{c}\right) & =\frac{\omega_{c}}{\gamma_{c}-1} \\
\chi^{\prime}\left(\gamma_{c}\right) & =\frac{\chi\left(\gamma_{c}\right)}{\gamma_{c}-1} \tag{119}
\end{align*}
$$

where the second equality follows from evaluation of (114) at $\gamma_{c}$. (119) can also be rewritten using the symmetry of $\chi(\gamma)$ in its principle branch: $\chi(1-\gamma)=\chi(\gamma)$ and $\chi^{\prime}(1-\gamma)=-\chi^{\prime}(\gamma)$. Letting $1-\gamma_{c} \rightarrow \gamma_{c}$,

$$
\begin{equation*}
\chi^{\prime}\left(\gamma_{c}\right)=\frac{\chi\left(\gamma_{c}\right)}{\gamma_{c}} \tag{120}
\end{equation*}
$$

This matching condition was actually first derived in the extensive 1983 Gribov, Levin, and Ryskin paper [48], but was rederived by Levin and Bartels in 1992 [42] with a more modern presentation.
More recently, in 2003 Munier and Peschanski [14] discovered a satisfying physical interpretation of the long known condition. Solving the linear part of the BK equation (97) as a wave packet in Mellin space,

$$
\begin{equation*}
N(k, Y)=\int_{c-i \infty}^{c+i \infty} \frac{d \gamma}{2 \pi i} N_{0}(\gamma) e^{-\gamma L+\bar{\alpha} \chi(\gamma) Y} \tag{121}
\end{equation*}
$$

we see that the phase velocity of a wave is

$$
\begin{equation*}
v_{p}=\frac{\chi(\gamma)}{\gamma} \tag{122}
\end{equation*}
$$

and the group velocity is

$$
\begin{equation*}
v_{g}=\frac{d \chi(\gamma)}{d \gamma} \tag{123}
\end{equation*}
$$

For the initial conditions relevant in QCD (a steeply falling function of L), FKPP analysis shows that the group velocity will equal the minimum phase velocity, which occurs at $\gamma=\gamma_{c}$.

$$
\begin{gather*}
v_{g}=\left.v_{p}\right|_{\min }=\frac{\chi\left(\gamma_{c}\right)}{\gamma_{c}}  \tag{124}\\
\chi^{\prime}\left(\gamma_{c}\right)=\frac{\chi\left(\gamma_{c}\right)}{\gamma_{c}} \tag{125}
\end{gather*}
$$

which is the same as 120 .
Before continuing, we will briefly address the nondeterministic nature of the evolution of the saturation scale. All that we have thus far discussed is deterministic and applies only to the mean field. However, because the formation of discrete dipoles ahead of the saturation front is a stochastic process, there will be some inherent dispersion among different "events", or realizations of BK evolution. As of currently, there has not been a rigorous proof of the behavior of this dispersion, but several numerical implementations have shown that

$$
\begin{equation*}
\sigma^{2} \propto Y \tag{126}
\end{equation*}
$$

There has been some progress in establishing this behavior using a "phenomenological" approach (see [18, 19]).

## Part II

## Model

## 5 Description of the Model: 2D, 2DR, and 2DSR

### 5.1 Overview

The object of our model is to implement Mueller's 2D branching kernel using a computer simulated Monte Carlo dipole generator. We expect the results to reproduce broad features of the FKPP traveling wave solution, in particular that the amplitude will behave like in figure 2 that we showed in the introduction, traveling with a fixed asymptotic velocity. Part of the motivation for this undertaking is to evaluate the following statement.

Note that, though a full study with two transverse degrees of freedom would be of great interest, we believe that our one-dimensional picture grasps the important aspects of the problem and, based on universal properties of the reaction-diffusion systems, we expect our results to hold for full QCD. 24]

Will a 2D model reproduce the same universal properties as the 1D model? In what ways will the details be refined? We seek to answer these questions.

First let us define a model "event". An event begins with an initial set of dipoles of size $r_{0}=1$ randomly oriented and randomly distributed in impact parameter such that $|\mathbf{b}|<\frac{r_{0}}{2}$. Over the course of evolution in time ${ }^{8}$, this initial dipole will have evolved into a multitude of smaller dipoles in each size index, exponentially at first but then tamed by a saturation mechanism. Each event consists of the movement of the saturation front $\rho_{s}$ to successively smaller sizes over a specified time interval. Because we expect the solution to take the form of a traveling wave, the amplitude should be a function only of

$$
\begin{equation*}
T\left(\rho-\rho_{s}(Y)\right)=T\left(\frac{k^{2}}{Q_{s}^{2}(Y)}\right) \tag{127}
\end{equation*}
$$

[^6]

Figure 15: Geometry of a dipole-dipole scattering.
if $\rho \sim 1 / r \sim k$. Thus, we see that $\rho_{s}(Y)$ plays the role of the saturation scale in the problem, and the traveling wave solution is equivalent to geometric scaling.
The amplitude can be calculated by making use of the following equation [17, 1].

$$
\begin{equation*}
T\left(y, \mathbf{x}_{01}\right)=\int \frac{d^{2} z_{0}}{2 \pi} \frac{d^{2} z_{1}}{2 \pi} T^{e l}\left(\mathbf{x}_{01}, \mathbf{z}_{01}\right) n\left(y, \mathbf{z}_{01}\right) \tag{128}
\end{equation*}
$$

where $n\left(y, \mathbf{z}_{01}\right)$ is the dipole density, and the elementary scattering amplitude for a projectile dipole scattering off a target dipole is

$$
\begin{equation*}
T^{e l}\left(\mathbf{x}_{01}, \mathbf{z}_{01}\right)=\frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2} \frac{\left|\mathbf{x}_{0}-\mathbf{z}_{1}\right|^{2}\left|\mathbf{x}_{1}-\mathbf{z}_{0}\right|^{2}}{\left|\mathbf{x}_{0}-\mathbf{z}_{0}\right|^{2}\left|\mathbf{x}_{1}-\mathbf{z}_{1}\right|^{2}} \tag{129}
\end{equation*}
$$

This formula represents the exchange of two gluons between a pair of dipoles, and as such is the square of the the single gluon potential between two dipoles in two dimensions [22]. It roughly counts the number of dipoles of similar size to $x_{01}$, which is convenient for computer implementation. Let us evince this feature. Given two dipoles of size $2 r$ and $2 R$, using the points shown on figure $15 T^{e l}$ can be written

$$
\begin{equation*}
T^{e l}=\frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2} \frac{\left(A B^{\prime}\right)^{2}\left(A^{\prime} B\right)^{2}}{(A B)^{2}\left(A^{\prime} B^{\prime}\right)^{2}} \tag{130}
\end{equation*}
$$

Case 1: $b \gg r, R$, leading order in $R^{2} / b^{2}, r R / b^{2}$, and $r^{2} / b^{2}$ :

$$
\begin{align*}
T^{e l} & =\frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2} \frac{\left[b^{2}+(R+r)^{2}\right]^{2}}{\left[b^{2}+(R-r)^{2}\right]^{2}} \\
& \approx \frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2}\left\{\left[1+2 \frac{(R+r)^{2}}{b^{2}}\right]\left[1-2 \frac{(R-r)^{2}}{b^{2}}\right]\right\} \\
& \approx \frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2}\left(1+\frac{8 r R}{b^{2}}\right) \\
& \approx \frac{\pi^{2} \alpha_{s}^{2}}{2}\left(\frac{8 r R}{b^{2}}\right)^{2}=32 \pi^{2} \alpha_{s}^{2} \frac{(r R)^{2}}{b^{4}} \sim \frac{(r R)^{2}}{b^{4}} \tag{131}
\end{align*}
$$

Case 2: $R>r, b$, leading order in $r / R$ and $b / R$ :

$$
\begin{align*}
T^{e l} & =\frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2} \frac{\left[b^{2}+(R+r)^{2}\right]^{2}}{\left[b^{2}+(R-r)^{2}\right]^{2}} \\
& \approx \frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2}\left[\left(1+\frac{4 r}{R}\right)\left(1+\frac{4 r}{R}\right)\right] \\
& \approx \frac{\pi^{2} \alpha_{s}^{2}}{2} \ln ^{2}\left(1+\frac{8 r}{R}\right) \\
& \approx \frac{\pi^{2} \alpha_{s}^{2}}{2}\left(\frac{8 r}{R}\right)^{2}=32 \pi^{2} \alpha_{s}^{2} \frac{r^{2}}{R^{2}} \sim \frac{r^{2}}{R^{2}} \tag{132}
\end{align*}
$$

From (131) and (132), we see that dipoles which are far apart or which have very different sizes will not greatly contribute to (128).

### 5.2 Determination of splitting probabilities and lifetimes

Recall the transverse space kernel we derived in (30), which represents a classical branching probability ${ }^{9}$ :

$$
\begin{equation*}
\frac{d P_{x_{01} \rightarrow x_{02}, x_{12}}}{d Y}=\frac{x_{01}^{2}}{x_{12}^{2} x_{02}^{2}} \frac{d^{2} \mathbf{x}_{2}}{2 \pi} \tag{133}
\end{equation*}
$$

In order to derive an expression for the lifetime of a given size dipole and its probability of splitting into another size dipole, we will integrate (133) over $\mathbf{x}_{2}$. Changing coordinates to a polar coordinate system with origin $\mathbf{x}_{1}$ and expanding $x_{02}^{2}$ with the

[^7]law of cosines,
\[

$$
\begin{equation*}
\frac{d P_{x_{01}}}{d Y}=2 x_{01}^{2} \int_{0}^{2 \pi} \frac{d \phi}{2 \pi} \int_{r_{\min }}^{r_{\max }} \frac{d x_{12}}{x_{12}\left(x_{01}^{2}+x_{12}^{2}-2 x_{01} x_{12} \cos \phi\right)} \tag{134}
\end{equation*}
$$

\]

The lower limit $r_{\text {min }}$ on the radial integral cuts off the collinear singularity, as we did in (44), whereas the upper limit $r_{\max }$ exists for the sake of computer implementation, as will become clear below. The left diagram in figure 16 shows the integration region around the point $\mathbf{x}_{1}$, with radial integration performed in such a way as to capture the collinear singularity around this point. This diagram depicts the parent dipole $x_{01}$ splitting into two daughter dipoles, $x_{12}$ and $x_{02}$. The placement of $\mathbf{x}_{2}$ determines both the lengths and positions of said daughters. Impact parameters $\left(b_{01}, b_{02}, b_{12}\right)$ are defined to be the midpoint of the line segment joining the two endpoints of a given dipole. The result of this particular process will be two daughter dipoles with the parent removed.

Although it might be tempting to extend the integration region to the entire plane in such a polar coordinate system, there are two problems associated with doing so. First, using the logarithmic indexing shown in figure 16 left (which will be defined shortly), notice that if $x_{12}=x_{01}$ and if $\phi=0$, measured with respect to the axis defined by $x_{01}$, then $x_{02}=0$ and the integrand in blows up. Of course, one could rotate the polar coordinate grid off of the singularity, but this brings us to our second point: symmetry dictates that we include the collinear singularity at $\mathbf{x}_{0}$ as well as $\mathbf{x}_{1}$. A simple method for doing so is to restrict the integration region to the vicinity of $\mathbf{x}_{1}$ and multiply by 2 to account for the symmetric probability distribution around $\mathbf{x}_{0}$. This accounts for the factor of 2 in (134).

So far we have only discussed how to capture the collinear singularity, but we must also include the infrared singularity when $x_{02}, x_{12} \gg x_{01}$ for our model to contain the proposed physics. Figure 16 right shows a scheme for covering most of the plane without overlap between the $\mathbf{x}_{0}$ and $\mathbf{x}_{1}$ regions. In practice we will divide the azimuthal range into 12 bins. Splittings of $x_{01}$ to equal size daughter $x_{12}$ are allowed in the azimuthal range $\frac{\pi}{3} \leq \phi<\frac{5 \pi}{3}$, shaded in green, while all splittings to larger sizes are restricted to $\frac{\pi}{2} \leq \phi<\frac{3 \pi}{2}$, shaded in yellow.
Continuing with the integral in (134) but switching to variable limits on $\phi$,


Figure 16: "Dartboard" diagrams indicating integration regions in (134). Left: Parent dipole $x_{01}$ splitting into daughter dipoles $x_{02}$ and $x_{12}$. The integration region is shown in the vicinity of $\mathbf{x}_{1}$. Right: The collinear region from the left figure is shaded in red, the equal size splitting region in green, and the infrared region in yellow. Only the first larger size splitting is shown for the infrared region, but the yellow region is understood to be an infinite radius section of a semicircle. The union of these three regions is mirrored for the region around $\mathbf{x}_{0}$.

$$
\begin{align*}
& \frac{d P_{x_{01}}}{d Y}=\frac{1}{\pi} \int_{\phi_{1}}^{\phi_{2}} d \phi \int_{r_{\text {min }}}^{r_{\text {max }}} \frac{d x_{12}}{x_{12}\left(1+\frac{x_{12}^{2}}{x_{01}^{2}}-2 \frac{x_{12}}{x_{01}} \cos \phi\right)}  \tag{135}\\
= & \frac{1}{\pi} \ln (B) \int_{\phi_{1}}^{\phi_{2}} d \phi \int_{\rho_{\min }}^{\rho_{\max }} \frac{d \rho}{1+B^{-2\left(\rho-\rho_{x}\right)}-2 B^{-\left(\rho-\rho_{x}\right)} \cos \phi} \tag{136}
\end{align*}
$$

where logarithmic sizes are defined by $\rho:=\log _{B}\left(\frac{1}{x_{12}}\right)$ and $\rho_{x}:=\log _{B}\left(\frac{1}{x_{01}}\right)$. The base $B$ determines the coarseness of the graining and will be taken to be 2 in the computer implementation of the model. Also, let $\rho_{\min }:=\log _{B} \frac{1}{r_{\max }}=0$ and $\rho_{\max }:=$ $\log _{B} \frac{1}{r_{\text {min }}}=50$ comprise the size limits on dipoles in our mode ${ }^{10}$. We will approximate this integral as a Riemann sum for the purposes of computer implementation, with $\Delta \phi$ and $\Delta \rho$ chosen to be, respectively, $\frac{2 \pi}{n}$ and 1. For $\rho_{\min } \leq \rho \leq \rho_{x}$ the angular region will be restricted, as discussed above.

$$
\begin{align*}
\frac{d P_{x_{01}}}{d Y} \approx & \frac{1}{\pi} \ln (B) \sum_{\rho=\rho_{\min }}^{\rho_{x}} \sum_{k=k_{1}}^{k_{2}} \frac{2 \pi}{n} \frac{1}{1+B^{-2\left(\rho-\rho_{x}\right)}-2 B^{-\left(\rho-\rho_{x}\right)} \cos \phi_{k}} \\
& +\frac{1}{\pi} \ln (B) \sum_{\rho=\rho_{y}+1}^{\rho_{\max }^{-1}} \sum_{k=0}^{n-1} \frac{2 \pi}{n} \frac{1}{1+B^{-2\left(\rho-\rho_{x}\right)}-2 B^{-\left(\rho-\rho_{x}\right)} \cos \phi_{k}} \tag{137}
\end{align*}
$$

Letting $i=\rho_{x}$ and $j=\rho$,

$$
\begin{equation*}
=\left.\sum_{j=\rho_{\min }}^{i} \frac{d P_{i \rightarrow j}}{d Y}\right|_{j \leq i}+\left.\sum_{j=i+1}^{\rho_{\max }-1} \frac{d P_{i \rightarrow j}}{d Y}\right|_{j>i} \tag{138}
\end{equation*}
$$

since

$$
\begin{aligned}
& \left.\frac{d P_{i \rightarrow j}}{d Y}\right|_{j>i}=\frac{1}{\pi} \ln (B) \int_{0}^{2 \pi} d \phi \int_{j}^{j+1} \frac{d \rho}{1+B^{-2(\rho-i)}-2 B^{-(\rho-i)} \cos \phi} \\
& \quad=\frac{1}{\pi} \ln (B) \sum_{\rho=j}^{(j+1)-1} \sum_{k=0}^{n-1} \frac{2 \pi}{n} \frac{1}{1+B^{-2(\rho-i)}-2 B^{-(\rho-i)} \cos \phi_{k}}
\end{aligned}
$$

[^8]\[

$$
\begin{equation*}
=\frac{1}{\pi} \ln (B) \sum_{k=0}^{n-1} \frac{2 \pi}{n} \frac{1}{1+B^{-2(j-i)}-2 B^{-(j-i)} \cos \phi_{k}} \tag{139}
\end{equation*}
$$

\]

and likewise,

$$
\begin{equation*}
\left.\frac{d P_{i \rightarrow j}}{d Y}\right|_{j \leq i}=\frac{1}{\pi} \ln (B) \sum_{k=k_{1}}^{k_{2}} \frac{2 \pi}{n} \frac{1}{1+B^{-2(j-i)}-2 B^{-(j-i)} \cos \phi_{k}} \tag{140}
\end{equation*}
$$

Thus, according to (138), the total probability for a dipole to split is the sum of the probabilities for it to split to any other size. For convenience, let us now define a probability splitting matrix $\mathcal{P}$ such that $\mathcal{P}_{i j k}$ is the $k$ th term in the azimuthal sum of $\frac{d P_{i \rightarrow j}}{d Y}$, i.e.

$$
\begin{equation*}
\mathcal{P}_{i j k}:=\frac{1}{\pi} \ln (B) \frac{2 \pi}{n} \frac{1}{1+B^{-2(j-i)}-2 B^{-(j-i)} \cos \phi_{k}} \tag{141}
\end{equation*}
$$

The $\mathcal{P}_{i j k}$ terms for which $\phi_{k}$ lies outside the azimuthal boundaries shown in figure 16 are set to 0 . We can now write the total probability for the splitting of $x_{01}$ (logarithmic size $i$ ) as

$$
\begin{equation*}
\frac{d P_{i}}{d Y}=\sum_{j=\rho_{\min }}^{\rho_{\max }-1} \sum_{k=0}^{n-1} \mathcal{P}_{i j k} \tag{142}
\end{equation*}
$$

and therefore, its "lifetime" in units of rapidity is

$$
\begin{equation*}
\tau_{i}=\left(d P_{i} / d Y\right)^{-1} \tag{143}
\end{equation*}
$$

The preceding forms the basis of our Monte Carlo calculation. During each step of the target's evolution in rapidity, the number of splittings of size $i$ is determined according to

$$
\begin{equation*}
\# \text { splittings }_{i}=\frac{1}{\tau_{i}} \Delta Y \times(\# \text { dipoles of size i }) \tag{144}
\end{equation*}
$$

We then randomly select this number of dipoles of size $i$, and for each selection choose a size $j$ to split into using the discrete probability distribution

$$
\begin{equation*}
\frac{d P_{i \rightarrow j}}{d Y}=\sum_{k=0}^{n-1} \mathcal{P}_{i j k} \tag{145}
\end{equation*}
$$

This can be done, for example, by randomly choosing a number on the interval $[0,1]$ in the properly normalized cumulative distribution function of (145) and finding the corresponding ordinate. Similarly, we can randomly choose an azimuthal bin $k$ to split into using the discrete probability distribution $\mathcal{P}_{i j k}$ for a given $i$ and $\sqrt{11}$.

### 5.3 Determination of $\mathrm{x}_{2}$

Once we have determined to which $j$ and $k$ a given dipole $x_{01}$ will split, it is a simple matter to locate $\mathbf{x}_{2}$. If splitting from $\mathbf{x}_{1}$,

$$
\begin{gather*}
\mathbf{x}_{2}=\mathbf{x}_{1}-r_{j} \mathcal{R}\left(\frac{2 \pi k}{n}\right) \hat{\mathbf{x}}_{01}  \tag{146}\\
\mathbf{x}_{2}=\mathbf{x}_{1}-r_{j} \mathcal{R}\left(\frac{2 \pi k}{n}\right) \frac{\mathbf{x}_{1}-\mathbf{x}_{0}}{\left|\mathbf{x}_{1}-\mathbf{x}_{0}\right|} \tag{147}
\end{gather*}
$$

where $\mathcal{R}(\theta)$ is the standard rotation matrix,

$$
\mathcal{R}(\theta):=\left(\begin{array}{cc}
\cos \theta & -\sin \theta  \tag{148}\\
\sin \theta & \cos \theta
\end{array}\right)
$$

By components,

$$
\begin{align*}
& x_{2 x}=x_{1 x}-r_{j}\left(\cos \theta \hat{x}_{01, x}-\sin \theta \hat{x}_{01, y}\right) \\
& x_{2 y}=x_{1 y}-r_{j}\left(\sin \theta \hat{x}_{01, x}+\cos \theta \hat{x}_{01, y}\right) \tag{149}
\end{align*}
$$

and

$$
\begin{equation*}
\mathbf{b}_{02}=\frac{\mathbf{x}_{0}+\mathbf{x}_{2}}{2}, \quad \mathbf{b}_{12}=\frac{\mathbf{x}_{1}+\mathbf{x}_{2}}{2} \tag{150}
\end{equation*}
$$

[^9]If splitting from the $\mathbf{x}_{0}$ side, then 146 becomes

$$
\begin{equation*}
\mathbf{x}_{2}=\mathbf{x}_{0}+r_{j} \mathcal{R}\left(\frac{2 \pi k}{n}\right) \hat{\mathbf{x}}_{01} \tag{151}
\end{equation*}
$$

mutatis mutandis.

### 5.4 Saturation veto and impact parameter cutoff veto

Limiting the number of dipoles in our model serves the dual purpose of satisfying unitarity constraints and ensuring computational efficiency. Toward this end, we will introduce two types of splitting vetoes into our model: the saturation veto and the impact parameter cutoff veto.

The former is based on the well known effect resulting from the BK equation, as discussed in 4.1. While the exact mechanism for saturation is not precisely known, be it a gluon recombination or shadowing effect, the results of our simulation should not strongly depend on the details. We will use the same condition as in [24, 25], which is that splittings that would generate daughters in regions already containing more than some $N_{\text {sat }}$ number of dipoles will not be allowed. But how are we to count the number of such dipoles?
Observing figure 17, say we want to probe the number of dipoles of logarithmic size $i$ in the vicinity of some impact parameter $\mathbf{b}_{p}$. We will count the number of dipoles whose impact parameters lie within an open ball around $\mathbf{b}_{p}, B_{r_{i} / 2}\left(\mathbf{b}_{p}\right):=$ $\left.\left\{\mathbf{b} \in \mathbb{R}^{2} \mid d\left(\mathbf{b}, \mathbf{b}_{p}\right)<r_{i} / 2\right\}\right\}^{12}$. Thus, in the figure the dipole with impact parameter $\mathbf{b}_{1}$ (shaded blue) is counted while that with $\mathbf{b}_{2}$ (shaded green) is not. However, even if this number of counted dipoles is less than $N_{\text {sat }}$, this does not guarantee that the saturation condition is not violated elsewhere. For example, say there are already $N_{s a t}$ dipoles with impact parameters very near $\mathbf{b}_{2}$. The addition of a dipole with impact parameter $\mathbf{b}_{p}$ will violate the saturation condition at some $\mathbf{b}_{3} \in B_{r_{i} / 2}\left(\mathbf{b}_{p}\right) \cup B_{r_{i} / 2}\left(\mathbf{b}_{2}\right)$, even though fewer than $N_{s a t}$ dipoles have impact parameters within $B_{r_{i} / 2}\left(\mathbf{b}_{p}\right)$. Thus, technically speaking we should check saturation at all $\mathbf{b} \in B_{r_{i} / 2}\left(\mathbf{b}_{p}\right)$ to ensure the saturation condition is never violated, but in practice saturation checks are very computationally expensive to carry out. Our results show that if checks are carried out at $\mathbf{b}_{p}, \mathbf{b}_{p}-\frac{r_{i}}{2} \hat{\mathbf{b}}_{p}$, and $\mathbf{b}_{p}+\frac{r_{i}}{2} \hat{\mathbf{b}}_{p}$, amplitudes obey saturation, and these checks are

[^10]

Figure 17: Two dipoles are shown with impact parameters $\mathbf{b}$ satisfying $\left|\mathbf{b}_{p}\right|-r_{i} / 2<$ $|\mathbf{b}|<\left|\mathbf{b}_{p}\right|+r_{i} / 2$. The dipole with impact parameter $\mathbf{b}_{1}$ (blue) is counted as being in the vicinity of $\mathbf{b}_{p}$ while that with $\mathbf{b}_{2}$ (green) is not. The crosshatched annulus is relevant to our search algorithm explained in 5.5.
ipso facto sufficient.
The other type of veto, which is a distance cutoff, is very easy to implement and necessary for computation in any reasonable length of time. If we choose a particular impact parameter $\mathbf{b}_{p}$ or set of impact parameters $\left\{\mathbf{b}_{p 1}, \mathbf{b}_{p 2} \ldots\right\}$ at which to check the amplitude throughout the evolution of an event, most dipoles-especially very small sizes-will be too far from any of the $\mathbf{b}_{p}$ for them or their progeny to affect $T\left(\mathbf{b}_{p}\right)$. Therefore, we impose the same cutoff as in 24,

$$
\begin{equation*}
\frac{r_{i}}{\left|\mathbf{b}-\mathbf{b}_{p n}\right|}>\kappa \tag{152}
\end{equation*}
$$

for some chosen value of $\kappa$ in order to allow the splitting which creates a daughter dipole at $\mathbf{b}$ with size $r_{i}$. 152 must be satisfied for at least one of the $\left\{\mathbf{b}_{p n}\right\}$ for the splitting to be allowed; otherwise it is vetoed. We can see that this condition results in smaller dipoles being more strongly constrained to the probe location(s):

$$
\begin{equation*}
\left|\mathbf{b}-\mathbf{b}_{p n}\right|<\frac{r_{i}}{\kappa} \tag{153}
\end{equation*}
$$

which is desirable, as there is no reason to keep track of the profusion of small dipoles that will not be observed. Typical values of $\kappa$ we will be using are $10^{-1}$ and $10^{-2}$. As long as $\kappa$ is not close to 1 , the asymptotic results of our model will not be greatly affected.

### 5.5 Data structure

2D evolution is much more computationally intensive than 1D due to the fact that a 2D transverse space can accommodate a far larger number of dipoles. Even given the veto constraints above, we must thoughtfully construct our data structure for computational efficiency. We can easily estimate the number of dipoles allowed for a given size $i$ using (153). Say $\mathbf{b}_{p}=\mathbf{0}$, then dipoles of size $i$ are constrained to a disk of radius

$$
\begin{equation*}
b_{i}<\frac{r_{i}}{\kappa} \tag{154}
\end{equation*}
$$

The number of dipoles that can exist within this radius is approximately


Figure 18: The data structure used to store dipoles. It is a vector with $\rho_{\max }+1$ entries, each of which a red-black tree header node. Each red-black tree is ordered by magnitude of impact parameter.

$$
N_{i} \approx N_{s a t} \frac{\pi b_{i}^{2}}{\pi\left(\frac{r_{i}}{2}\right)^{2}}=\frac{4 N_{s a t}}{\kappa^{2}}
$$

For typical values we will be using, $\kappa=10^{-1}$ and $N_{s a t}=25, N_{i} \approx 10,000$. We have discovered that a 2D simulation becomes very computationally unwieldy when $N_{i} \gtrsim 10^{5}$. For this reason, $\kappa=10^{-1}$ will be our standard choice for full 2D simulation. The main data structure of the program will contain all of the dipoles created in the course of the target's evolution. It will consist of a vector $\left\{n_{i}\right\}_{i \in\left\{0,1,2, \ldots, \rho_{\max }\right\}}$, each index $i$ of which represents all dipoles of logarithmic size $i$. The vector object type will be a binary red-black tree of nodes ordered by magnitude of impact parameter and that each contain the variables $\left\{b, b_{x}, b_{y}, x_{0 x}, x_{0 y}, x_{1 x}, x_{1 y}\right\}$. This is indicated schematically in figure 18.

Let us divert our attention to the red-black tree structure for each size index, which is crucial to the program's ability to quickly carry out saturation checks of the type described in 5.4. The conceptual basis for the red-black tree can be found in a number of references, for example its inventor's textbook, [50], but we will summarize the basic features here for the reader less familiar with data structures. Essentially, the redblack tree's purpose is to maintain the binary search tree's (BST) optimal $O\left(\log _{2} N\right)$ search performance. It is one of several self-balancing tree algorithms available ${ }^{[13}$.

[^11]

Figure 19: Left: A low efficiency BST with $O(N)$ search time. Right: A high efficiency BST with $O\left(\log _{2} N\right)$ search time.

Consider the degenerate case of adding, in sequence, $1,2,3,4,5$ to a standard BST (figure 19). The insertion algorithm for a BST is to traverse the tree, going left if the node to be inserted is smaller than the current tree node, and right if it is greater. Thus, figure 19 left obtains with search time $O(N)$, as the BST degenerates into essentially a linked list in such cases. Figure 19 right obtains if we insert the sequence $2,1,4,3,5$, but we would like to achieve this efficient $O\left(\log _{2} N\right)$ structure independent of insertion order. That is where the red-black tree comes into play.
A red-black tree's insertion and deletion algorithms ensure that its branches will remain roughly balanced at all times by leaving the following properties intact:

1. Each node is either red or black.
2. The root node is black.
3. Both children of every red node are black. If unsatisfied, there is said to be a "red violation".
4. Every path from root to leaf ${ }^{14}$ contains the same number of black nodes. If unsatisfied, there is said to be a "black violation".

Such a tree satisfying these properties is shown in figure 20, as the reader may verify. Although the rebalancing algorithms are fairly detailed and refer to a number of

[^12]

Figure 20: A sample red-black tree, ordered by magnitude of impact parameter
different cases, we will give one example to indicate the flavor of the operations required.

Say we are adding the node with impact parameter value ".93664". The red-black tree will now look like figure 21 upper. We can see that there is currently a red violation since the new node and its parent are both red. We cannot simply recolor the new node black, as this would lead to a black violation. Instead, we will recolor the new node's parent and grandparent, as shown in the diagram. Unfortunately, this causes another red violation. We cannot again recolor grandparent and great grandparent, as this would violate property 2 . Thus, we can see that rotations are required for rebalancing. These rotations, along with recoloration and reattachment of appropriate subtrees are indicated in figure 21 middle. We end up with figure 21 bottom, which has the immediate visual appearance of being more balanced than 21 top.

Without going through all of the cases, suffice it to say that algorithms exist to maintain properties 1 through 4 during insertion and deletion of nodes. (The latter is especially tedious and is usually omitted from texts.) Several different types of algorithms actually exist to accomplish these tasks. The example given above is a type of "bottom-up" algorithm which recursively travels up the tree from the insertion point fixing mistakes on the way up. Another method involves nodes which have pointers from children to parents as well as from parents to children. However, both of these


Figure 21: An example of red-black tree rebalancing after adding the node containing ". 93664 " on the far right.
methods appear somewhat inelegant when compared with "top-down" insertion. Topdown insertion is a nonrecursive method that makes changes on the way down the tree to the insertion point. Since it does useful work on the way down and does not have to traverse back up the tree, it is the most efficient method of implementing the red-black tree. It is surprisingly difficult to find these algorithms, but 51] provides a discussion of them.

Having the red-black tree data structure at our disposal allows us to quickly check $\left\{n_{i}\right\}$ for saturation vetoes, as explained in 5.4, and also to calculate the $T_{i}\left(\mathbf{b}_{p}\right)$, the amplitude at $\mathbf{b}_{p}$ for size $i$ dipoles, at each step of the target's evolution. This is done by searching $n_{i}$, the $i$ th red-black tree, for dipoles satisfying

$$
\begin{equation*}
\max \left(0, b_{p}-\frac{r_{i}}{2}\right)<b<b_{p}+\frac{r_{i}}{2} \tag{155}
\end{equation*}
$$

This check is efficiently accomplished given the $O\left(\log _{2} N\right)$ search performance of the red-black tree. Notice that (155) corresponds to the annulus in figure 17. Of course, we also need to check each dipole satisfying (155) to see whether

$$
\begin{equation*}
\left|\mathbf{b}_{p}-\mathbf{b}\right|<\frac{r_{i}}{2} \tag{156}
\end{equation*}
$$

which is the number of dipoles with impact parameters within the open ball $B_{r_{i} / 2}\left(\mathbf{b}_{p}\right)$, shaded red in figure 17. The sum of dipoles that satisfy (155) and (156) divided by $N_{\text {sat }}$ yields $T_{i}\left(\mathbf{b}_{p}\right)$. Knowledge of $T_{i}\left(\mathbf{b}_{p}\right)$ for all $i$ also allows us to calculate the saturation front, $\rho_{s}\left(\mathbf{b}_{p}, Y\right)$, which we will define as the smallest $i$ such that $T_{i}\left(\mathbf{b}_{p}\right)<\frac{1}{2}$.

### 5.6 Parallel coding

Our C++ code was written using the OpenMP API for shared memory multiprocessing. Threading is controlled by the use of "\#pragma" directives in the code, which stands for "pragmatic". These allow the $\mathrm{C}++$ compiler to precisely control memory management and passing of parameters so as to offer machine and operating systemspecific features while maintaining $\mathrm{C}++$ compatibility. This platform independence allows the programmer to run the same code on machines of different number of cores while always utilizing the maximum advantage of multithreading on each machine.

Short data runs were performed on a typical home PC with 4 cores running at 2.67 GHz while longer runs up to 24 hours were performed on the Texas Advanced Com-

| System Name: | Lonestar 4 |
| :---: | :---: |
| Host Name: | lonestar.tacc.utexas.edu |
| Operating System: | Linux |
| Number of Processors: | 22,656 |
| Total Memory | 44 TB |
| Peak Performance | 302 TFLOPS |
| Total Disk: | $276 \mathrm{~TB}($ local $), 1000 \mathrm{~TB}$ (global) |

Table 2: TACC Lonestar 4 specifications
puter Center's (TACC) Lonestar 4 Dell Linux cluster. Without going into great detail, the basic specifications of this cluster are the following: [52]

The 22,656 cores are housed on 1,888 Dell PowerEdge M610 compute blades with 12 to a blade. Each blade has 2 Xeon 5680 series 3.33 GHz hex-core processors. The user may submit jobs serially to each compute blade, which then multithreads the code onto 12 cores, providing essentially a 12 -fold increase in the rate data production for our simulation. Multiple blades may be simultaneously harnessed, allowing further generation of data.

### 5.7 Pseudocode program

Most of what has not been described heretofore is merely nuts and bolts of programming, such as declarations, flow control statements, data output, and the like. The essential physics has all been described. For the reader interested in how the program works, we will give a pseudocode overview of the program flow. This description is for a single event-multiple events are simply repeated instances of a single event.

## Program flow, single event

- main rapidity loop over $Y$ :
- loop over dipole size $i$ :
* calculate number of splittings for size $i$ using lifetime, see (144)
* loop over number of splittings, $l$ :
- choose a random dipole of size $i$ to split
- monte carlo this dipole into size $j$ dipole, see 145
- monte carlo into $k$ th angular bin
- randomly choose which side of dipole $i$ to split, see (146) and (151)
- check if 2 daughter dipoles, $\mathbf{x}_{02}$ and $\mathbf{x}_{12}$, satisfy $\kappa$ cutoff (152); if not, veto splitting
- check if 2 daughter dipoles, $\mathbf{x}_{02}$ and $\mathbf{x}_{12}$, violate saturation; if so, veto splitting
- if neither veto has been applied, insert $\mathbf{x}_{02}$ and $\mathbf{x}_{12}$ and remove $\mathbf{x}_{01}$ from the appropriate red-black trees in the data structure shown in figure 18
- output data for this $\Delta Y$ step


### 5.8 First several steps of an event

To illustrate the operations of the program, let us visually inspect the first several splittings of a single initial dipole. The program randomly generates the following two splittings during the first $\Delta Y$ step, as shown in figure 22 .

First Splitting:

$$
\begin{aligned}
& x_{0 x}=0.744071, x_{1 x}=-0.253879, x_{2 x}=-0.269879 \\
& x_{0 y}=-0.397142, x_{1 y}=-0.333142, x_{2 y}=-0.58263
\end{aligned}
$$

Second Splitting:

$$
\begin{aligned}
& x_{0 x}=-0.253879, x_{1 x}=-0.269879, x_{2 x}=-0.305895 \\
& x_{0 y}=-0.333142, x_{1 y}=-0.58263, x_{2 y}=-0.298492
\end{aligned}
$$

Both of these splittings occur in the collinear region, the first from $i=0$ to $j=2$ and the second from $i=2$ to $j=4$ (in logarithmic size). Notice that although the probability to split to a much smaller size is not improbable via 145), it is extremely improbable that a dipole created near the endpoints of its parent will pass the $\kappa$ cutoff condition (152), which is to say it will likely be too far away from the region of interest to have any effect there. Over the course of the evolution of an event, smaller size dipoles will have found their way sufficiently near the probe location via other


Figure 22: The first two splittings of an initial dipole shown in transverse space in clockwise progression. Removed parent dipoles are shown in red while extant dipoles are blue.


Figure 23: The evolution of a single dipole in transverse space at time $Y=1$.
somewhat larger sizes to pass (152). In this sense the traveling wave moves smoothly from larger to smaller size dipoles over the evolution time.

After a longer period of time, the parent dipole will have branched into a multitude of various sized smaller dipoles, shown in figure (23). These daughter dipoles remain a connected graph, as the splitting rules imply.
Beginning with $N_{\text {initial }}=N_{\text {sat }}=25$ dipoles and after sufficient time, a more fully evolved target is attained (figure 24).

### 5.9 2D Restricted (2DR)

It is desirable to have a way to check the results of our 2D calculation in the 1D limit in order to make contact with other work that has been done in 1D. To do so, we will employ the method illustrated in figure 25. The operation of the program is very similar to the 2D calculation, but with an added step before the veto conditions are checked. Recall that a logarithmic size and angle are chosen using discrete probability distributions, as described in 5.3. In the newly introduced step, the impact parameters $\mathbf{b}_{\mathbf{1 2}}$ and $\mathbf{b}_{02}$ are projected onto the $\mathbf{x}$-axis. If we were to simply project $\mathbf{x}_{2}$ onto the x -axis as well, this would have the effect of shortening the two projected dipoles $x_{12}$ and $x_{02}$, especially in the case of an infrared splitting. Instead, we want to preserve the lengths $x_{12}$ and $x_{02}$, which can be done by redefining the endpoints of the two daughters in the following way.
The endpoints of the dipole $x_{12}^{\prime}$ are given by


Figure 24: Going clockwise, target at time $Y=.5, Y=1, Y=1.5$, and $Y=2$, all with $N_{\text {initial }}=N_{\text {sat }}=25$ initial dipoles .


Figure 25: A method for reducing the full 2D calculation to 1D.

$$
\begin{align*}
& x_{1 x}^{\prime}=b_{12 x}+\frac{x_{12}}{2} \hat{x}_{01 x} \\
& x_{2 x}^{\prime}=b_{12 x}-\frac{x_{12}}{2} \hat{x}_{01 x} \tag{157}
\end{align*}
$$

and those of $x_{02}^{\prime}$ by

$$
\begin{align*}
x_{0 x}^{\prime \prime} & =b_{02 x}-\frac{x_{02}}{2} \hat{x}_{01 x} \\
x_{2 x}^{\prime \prime} & =b_{02 x}+\frac{x_{02}}{2} \hat{x}_{01 x} \tag{158}
\end{align*}
$$

In this scheme, we lose the shared endpoint between daughters, as $x_{2 x}^{\prime} \neq x_{2 x}^{\prime \prime}$, but dipole sizes of the 2D model are preserved.

### 5.10 2D Semi-Restricted (2DSR)

In another variation of our model, this time we would like to be able to smoothly transition from the full 2D calculation to a 1D version of that calculation. The basic idea is to allow dipoles to evolve by spreading in the azimuth, but only within a certain defined strip width $d$ around the x-axis. The shaded strip is shown in figure 26 left. Clearly the strip size must scale with the daughter dipole size if evolution is to be effectively constrained near the x-axis. We define $d$ in the following way:

$$
\begin{equation*}
d:=\beta r_{i} \tag{159}
\end{equation*}
$$

where $r_{i}=\min \left(x_{02}, x_{12}\right)$ is the size of the smaller daughter dipole and $\beta$ is a factor that mediates the transition from 2 D to 1 D . If $\mathbf{x}_{2}$ lies within the strip then no projection takes place. If, on the other hand $\left|x_{2 y}\right|>d$, then the projection

$$
\begin{equation*}
x_{2 y} \rightarrow x_{2 y}^{\prime}=s d, \quad 0<s<1 \tag{160}
\end{equation*}
$$

shown in 26, left takes place, with $s$ a random number in the interval above. In order to preserve the lengths of $x_{02}$ and $x_{12}$, we slide $\mathbf{x}_{0}$ and $\mathbf{x}_{1}$ along the x -axis away from


Figure 26: Left: Gluons emitted outside of a strip of width 2 d around the x -axis are projected into the strip (shaded yellow) a distance $s d$ away from the x-axis, where $0<s<1$. Right: The limit of 2DSR as the strip width $d \rightarrow 0$.
$x_{2 x}{ }^{15}$.

$$
\begin{gather*}
x_{02}^{\prime}=x_{02}, \quad x_{12}^{\prime}=x_{12} \\
x_{1 x}^{\prime}=x_{2 x}^{\prime} \pm \sqrt{x_{12}^{2}-\left(x_{1 y}-x_{2 y}^{\prime}\right)^{2}} \\
x_{0 x}^{\prime}=x_{2 x}^{\prime} \mp \sqrt{x_{02}^{2}-\left(x_{0 y}-x_{2 y}^{\prime}\right)^{2}} \tag{161}
\end{gather*}
$$

The end result of this scheme is that when $\beta \rightarrow \infty$ we recover the full 2 D calculation, and when $\beta \rightarrow 0$ the calculation becomes 1D, as shown in figure 26 right. Note that this 1 D limit is not exactly the same as the 2 DR scheme, although the differences in the overall results between the two are minor.

[^13]
## 6 Results and Analysis

### 6.1 2D results

In presenting our results, we will display a number of the following quantities. Recall that the saturation front $\rho_{s}(Y, b)$ is a function of $Y$ and $b$.

$$
\begin{gather*}
\frac{d \rho_{s}}{d Y}=\frac{\left\langle\rho_{s}(Y+\Delta Y, 0)-\rho_{s}(Y, 0)\right\rangle}{\Delta Y}  \tag{162}\\
\sigma^{2}=\left\langle\rho_{s}^{2}(Y, 0)\right\rangle-\left\langle\rho_{s}(Y, 0)\right\rangle^{2}  \tag{163}\\
\operatorname{Cov}(b):=\operatorname{Cov}\left(\rho_{s}(Y, 0), \rho_{s}(Y, b)\right)=\left\langle\rho_{s}(Y, 0) \rho_{s}(Y, b)\right\rangle-\left\langle\rho_{s}(Y, 0)\right\rangle\left\langle\rho_{s}(Y, b)\right\rangle \tag{164}
\end{gather*}
$$

And with these definitions,

$$
\begin{equation*}
\operatorname{Cov}(0)=\sigma^{2} \tag{165}
\end{equation*}
$$

as expected. Note that only ensemble averages are shown, and thus, individual events would have a more discrete appearance than the mean curves displayed on the amplitude plots. Also, individual events will be ahead of or behind the mean curves, the degree to which is indicated by the accompanying variance plots. Note that the attached $\mathrm{C}++$ code only outputs the amplitude at various impact parameters and times. Additional data processing was handled in Matlab.

Figure 27 reveals the asymptotic wave speed to be about 3.5 -much slower than the 1D models we will consider. Variance is proportional to $Y$ after an initial wavefront formation time, as we expect from (126). The explanation for the the saturated region in 27 left having an amplitude slightly higher than 1 is the fact that we have only performed saturation checks at three points in transverse space when adding dipoles, as explained in 5.4. However, this slight excess has little effect on asymptotic values.


Figure 27: 2D Model: 700 events, $\kappa=10^{-1}$

### 6.2 2DR results

Because the 2DR model is restricted to 1 D , the number of dipoles allowed is severely curtailed when compared to 2D (see beginning of 5.5 for details). It is thereby much easier to gather high statistics in this version of the dipole model. With 5000 events in figure 28, wavefront velocity and variance curves are the smoothest of the data we present. The three point saturation check is also clearly more effective in 1D, as amplitudes are kept below $T(Y)=1$ in the plots shown. Additionally, asymptotic wave velocity is seen to be much greater in 1D than in 2D, which we will discuss later. A comparison of figures 28 and 29 reveals that a change in $\kappa$ has little effect on asymptotic velocity: $\left\langle d \rho_{s} / d Y\right\rangle=14.078$ for the former while $\left\langle d \rho_{s} / d Y\right\rangle=14.390$ for the latter ${ }^{[16}$. It is slightly larger for the latter because $\kappa=30^{-1}$ for this data allows dipoles to form within a radius three times greater (at a given $i$ ) than $\kappa=10^{-1}$ for the former. Some of these additional dipoles that are farther from the probe location will be able to "walk in" through successive splittings. Further decreasing $\kappa$ will have a diminishing effect on the wavefront velocity since the farther away a dipole is from the probe, the less likely it is have an effect there.

Figure 30 displays decorrelation of wavefronts at various impact parameters. This phenomenon is intuitively explained by considering the "resolution" of dipoles required to distinguish between two points. As long as the dipoles present in the simulation are larger than the separation between two impact parameters, these impact parameters are correlated and their covariance will rise over time. The points will decorrelate

[^14]

Figure 28: 2DR Model: 5000 events, $\kappa=10^{-1}$



Figure 29: 2DR Model: 1000 events, $\kappa=30^{-1}$
(their covariance will become constant) when the event has reached a fine enough resolution such that [25]

$$
\begin{equation*}
\Delta b \approx B^{-\rho_{s}(Y)} \tag{166}
\end{equation*}
$$

Table 3 details the $Y$ values at which various impact parameters decorrelate from $b=0$. These $Y$ values match well with figure 30 .

### 6.3 2DSR results

The data from figure 31interpolates between the 1D and 2D realizations of our model. As $\beta$ increases, widening the projection strip, we see the essentially 1D results from the top row become the 2 D results from the last row.


Figure 30: 2DR Model: 300 events, $\kappa=10^{-1}$

| $\Delta b$ | $\rho_{s}$ | $Y_{\text {decor }}$ |
| :---: | :---: | :---: |
| $10^{-6}$ | 19.9 | 2 |
| $10^{-4}$ | 13.3 | 1.5 |
| $10^{-2}$ | 6.6 | 1 |
| $10^{-1}$ | 3.3 | .8 |

Table 3: Decorrelation data for impact parameters in figure 30 calculated using 166 .


Figure 31: 2DSR Model: 1st row: $\beta=0$; 2nd row: $\beta=1$; 3rd row: $\beta=3$; 4th row: $\beta=100$. All data 500 events and $\kappa=10^{-1}$.

### 6.4 Wavefront velocity analysis

### 6.4.1 1D Eigenvalue calculation

Splitting the kernel $K_{i j}$ into infrared $(j<i)$, collinear $(j>i)$, and equal size $(j=i)$ parts,

$$
\begin{align*}
\partial_{Y} n_{i} & =\sum_{j=\rho_{\min }}^{\rho_{\max }-1} K_{i j} n_{j} \\
& =\left.\sum_{j=\rho_{\min }}^{i} \frac{d P_{i \rightarrow j}}{d Y}\right|_{j<i} n_{j}+\left.\sum_{j=i+1}^{\rho_{\max }-1} \frac{d P_{i \rightarrow j}}{d Y}\right|_{j>i} n_{j}+\left.\frac{d P_{i \rightarrow j}}{d Y}\right|_{j=i} n_{j} \tag{167}
\end{align*}
$$

The splitting probability is given by the BFKL kernel, transformed to logarithmic size index:

$$
\begin{align*}
& \frac{d P_{i \rightarrow j}}{d Y} \equiv \frac{1}{\pi} \ln (B) \int_{0}^{2 \pi} d \phi \int_{j}^{j+1} \frac{d \rho}{1+B^{-2(\rho-i)}-2 B^{-(\rho-i)} \cos \phi} \\
&=\frac{1}{\pi} \ln (B) \sum_{\rho=j}^{(j+1)-1} \sum_{l=0}^{n-1} \frac{2 \pi}{n} \frac{1}{1+B^{-2(\rho-i)}-2 B^{-(\rho-i)} \cos \phi_{l}} \\
&=\frac{1}{\pi} \ln (B) \sum_{l=0}^{n-1} \frac{2 \pi}{n} \frac{1}{1+B^{-2(j-i)}-2 B^{-(j-i)} \cos \phi_{l}} \tag{168}
\end{align*}
$$

First we will handle the collinear term $(j>i)$. Using the following approximation with $\zeta:=\frac{x_{12}}{x_{01}}=B^{-(j-i)}$,

$$
\begin{align*}
\frac{1}{1+\zeta^{2}-2 \zeta \cos \phi} & =\sum_{m=0}^{\infty}\left(2 \zeta \cos \phi-\zeta^{2}\right)^{m} \\
& =1+\left(2 \zeta \cos \phi-\zeta^{2}\right)+2 \zeta^{2}(1+\cos 2 \phi)+O\left(\zeta^{3}\right) \\
& \rightarrow 1+\zeta^{2}+O\left(\zeta^{3}\right) \\
& \approx 1+B^{-2(j-i)} \tag{169}
\end{align*}
$$

where the identity $(2 \cos \phi)^{2}=2(1+\cos 2 \phi)$ was used in the second step, and the
integration of cosine terms set to 0 in the third, we can simplify the kernel. Using the eigenfunctions $\varphi_{j}=B^{j \gamma}$, and inserting a factor $\frac{r_{j}}{r_{i}}=B^{i-j}=: B^{-k}$ to reduce 1D to 0D fixed impact parameter (FIP),

$$
\begin{gather*}
\chi_{C O L, 1 D}(\gamma) \varphi_{i}(\gamma) \approx \sum_{j>i} \frac{1}{\pi} \ln (B) \sum_{l=0}^{n-1} \frac{2 \pi}{n} B^{-k}\left(1+B^{-2(j-i)}\right) \varphi_{j}(\gamma)  \tag{170}\\
=2 \ln (B)\left(\sum_{k>0} B^{-k} B^{\gamma k}\left(1+B^{-2 k}\right)\right) \varphi_{i}(\gamma) \\
\chi_{C O L, 1 D}(\gamma)=2 \ln (B)\left(\sum_{k>0} B^{k(\gamma-1)}+B^{k(\gamma-3)}\right) \\
=2 \ln (B)\left(\frac{1}{1-B^{\gamma-1}}+\frac{1}{1-B^{\gamma-3}}-2\right) \tag{171}
\end{gather*}
$$

Restoring the $\Delta$ factors, taking the limit as $\Delta \rightarrow 0$, and using L'Hospital's Rule,

$$
\begin{equation*}
\lim _{\Delta \rightarrow 0} \chi_{C O L}(\gamma)=\frac{2}{\gamma-1}+\frac{2}{\gamma-3} \tag{172}
\end{equation*}
$$

we see that the $\gamma=1$ singularity is present. Moving on to the infrared part $(j<i)$,

$$
\begin{equation*}
\chi_{I R, 1 D}(\gamma) \varphi_{i}(\gamma)=\left.\sum_{j=\rho_{\min }}^{i} B^{-k} \frac{d P_{i \rightarrow j}}{d Y}\right|_{j<i} \varphi_{j}(\gamma) \tag{173}
\end{equation*}
$$

Notice that since we are integrating semi-circles, we only sum over half of the azimuth.

$$
\begin{gather*}
=\sum_{k<0} \frac{1}{\pi} \ln (B) \sum_{l=0}^{\text {half }}{ }^{\text {azimuth }} \frac{2 \pi}{n} B^{-k} \frac{1}{1+B^{-2 k}-2 B^{-k} \cos \phi_{l}} B^{j \gamma} \\
\approx \ln (B) \sum_{k<0} B^{-k} B^{2 k} B^{j \gamma} \\
=\ln (B) \sum_{k<0} B^{k} B^{k \gamma} \varphi_{i}(\gamma) \\
\chi_{I R, 1 D}(\gamma)=\ln (B)\left(\frac{1}{1-B^{-(\gamma+1)}}-1\right) \tag{174}
\end{gather*}
$$

Finally, there is the $k=0$ term, whose integral bounds come from figure 16 ,

$$
\chi_{k=0,1 D}=\frac{1}{\pi} \ln (B) \int_{\pi / 3}^{5 \pi / 3} \frac{1}{1+B^{0}-2 B^{0} \cos \phi}=\frac{\sqrt{3}}{\pi} \ln (B)
$$

Adding the two parts (171) and (174),

$$
\begin{gather*}
\chi_{1 D}(\gamma)=\chi_{C O L, 1 D}(\gamma)+\chi_{I R, 1 D}(\gamma)+\chi_{k=0,1 D} \\
=2 \ln (B)\left(\frac{1}{1-B^{\gamma-1}}+\frac{1}{1-B^{\gamma-3}}+\frac{1}{2\left(1-B^{-(\gamma+1)}\right)}-\frac{5}{2}+\frac{\sqrt{3}}{2 \pi}\right) \tag{175}
\end{gather*}
$$

### 6.4.2 2D Eigenvalue calculation

Repeating all of the above steps but using instead the FIP factor of $\left(\frac{r_{j}}{r_{i}}\right)^{2}=B^{-2 k}$ to reduce 2D to 0D, (171) and (174) become

$$
\begin{gather*}
\chi_{C O L, 2 D}(\gamma)=2 \ln (B)\left(\frac{1}{1-B^{\gamma-2}}+\frac{1}{1-B^{\gamma-4}}-2\right)  \tag{176}\\
\chi_{I R, 2 D}(\gamma)=\ln (B)\left(\frac{1}{1-B^{-\gamma}}-1\right)  \tag{177}\\
\chi_{2 D}(\gamma)=2 \ln (B)\left(\frac{1}{1-B^{\gamma-2}}+\frac{1}{1-B^{\gamma-4}}+\frac{1}{2\left(1-B^{-\gamma}\right)}-\frac{5}{2}+\frac{\sqrt{3}}{2 \pi}\right) \tag{178}
\end{gather*}
$$

### 6.4.3 Velocity calculations

Notice that for neither 1D nor 2D do we get both poles. 171) has the $\frac{1}{\gamma-1}$ pole, and (177) has the $\frac{1}{\gamma}$ pole. This is perhaps to be expected since the 1D FIP correction factor $B^{-k}$ works well for the collinear sum in which dipoles remain more or less collinear. However, the 2D FIP factor $B^{-2 k}$ is better suited to the infrared sum since these kind of splittings allow the daughter dipoles to explore the azumithal range. We might consider using a "hybrid" eigenvalue function which has both of the correct poles,

$$
\chi_{h y b r i d}(\gamma) \equiv \chi_{C O L, 1 D}+\chi_{I R, 2 D}+\chi_{k=0}
$$

$$
\begin{equation*}
=2 \ln (B)\left(\frac{1}{1-B^{\gamma-1}}+\frac{1}{1-B^{\gamma-3}}+\frac{1}{2\left(1-B^{-\gamma}\right)}-\frac{5}{2}+\frac{\sqrt{3}}{2 \pi}\right) \tag{179}
\end{equation*}
$$

Using the eigenvalue functions (175), (178), and (179) and solving (120) using numerical methods, we obtain

$$
\begin{array}{cc}
V_{1 D}=\frac{\chi_{1 D}^{\prime}\left(\gamma_{c}\right)}{\ln (2)}=12.67, & \gamma_{c}=0.53 \\
V_{2 D}=\frac{\chi_{2 D}^{\prime}\left(\gamma_{c}\right)}{\ln (2)}=3.63, & \gamma_{c}=1.19 \\
V_{\text {hybrid }}=\frac{\chi_{\text {hybrid }}^{\prime}\left(\gamma_{c}\right)}{\ln (2)}=15.35 & \gamma_{c}=0.61 \tag{180}
\end{array}
$$

Comparing these values to the data, we see the our analytical calculation for $V_{2 D}$ looks very accurate. Using the data shown in figure 27 we obtain $\left\langle d \rho_{s} / d Y\right\rangle=3.513{ }^{177}$. The 2DR and 2DSR models suggest a value of $\left\langle d \rho_{s} / d Y\right\rangle=13.5$, which is still reasonably close to $V_{1 D}$. We can also calculate the asymptotic velocity from the actual BFKL eigenvalue function, $\chi(\gamma)=2 \psi(1)-\psi(1-\gamma)-\psi(\gamma)$. Using, for instance, [16]

$$
\begin{equation*}
\ln Q_{s}^{2}(Y)=\frac{\chi\left(\gamma_{c}\right)}{\gamma_{c}} Y-\frac{3}{2 \gamma_{c}} \ln (Y / \bar{\alpha})-\frac{3}{\gamma_{c}^{2}} \sqrt{\frac{2 \pi}{\chi^{\prime \prime}\left(\gamma_{c}\right)}} \frac{1}{\sqrt{Y}}+\mathcal{O}(1 / Y) \tag{181}
\end{equation*}
$$

the dominant term asymptotically yields

$$
\begin{align*}
\frac{d \ln Q_{s}^{2}(Y)}{d Y} & \approx \frac{\chi\left(\gamma_{c}\right)}{\gamma_{c}} \\
\frac{d \rho_{s}}{d Y} & =\frac{1}{2 \ln (2)} \chi^{\prime}\left(\gamma_{c}\right) \approx 3.523 \tag{182}
\end{align*}
$$

which also compares well with our 2D model value.

[^15]
### 6.5 Conclusions

One facet of wave propagation we have noticed is the necessity of including both the infrared and collinear singularities of the branching kernel. Because the saturation front propagates to smaller dipole sizes over time, the collinear part of the kernel drives the wave forward in $x$ while the infrared part "fills in" the unsaturated sizes behind the wavefront. Without the back-filling effect of the infrared term, the wave moves forward but is eventually damped out as the larger dipoles are replaced by dispersed smaller ones, and consequently, no stable wave shape asymptotically forms. Comparing 1D and 2D data, it is seen that average wavefront velocities are considerably higher for the former. We have not seen a discussion of this effect in previous work, probably because no previous work has undertaken a model in two dimensions. One explanation why the saturation front progresses faster in 1D configuration space than in 2D is that dipoles spreading out in 2D transverse space with the same splitting probability as used in the 1D model become more dilute in comparison. As long as daughter dipoles are confined to a line, it is much more probable that each splitting will increase dipole density near the probe than in 2 D . This reasoning still does not make the result a priori obvious, since one might imagine that the far more numerous dipoles in 2D could compensate for this dilution; however, it is seen that they do not. The analytical work in 6.4 gives some justification for this lower velocity.

We would like to consider the statement made in an earlier work,

Note that, though a full study with two transverse degrees of freedom would be of great interest, we believe that our one-dimensional picture grasps the important aspects of the problem and, based on universal properties of the reaction-diffusion systems, we expect our results to hold for full QCD. 24]

Let us take stock of some of the assumptions made in the [24] model:

- Parent dipoles are retained throughout the evolution; collinear splitting rules create one small daughter dipole while the maintained parent approximates the other daughter. Infrared splitting probabilities are increased by a factor of 2 since only one daughter is created-the parent is still maintained.
- Dipole size is discrete: all dipoles have a size $B^{-i}$ for some $i$.


Figure 32: Average wavefront velocity, as shown in [24]

- The 2D kernel 133 is replaced by a 1D version,

$$
\frac{d P}{d Y}=\frac{x_{01}}{x_{02} x_{12}} d x_{2}
$$

- The impact parameter of daughter dipoles is chosen using

$$
b_{j}=b_{i} \pm \frac{r_{i}}{2} \pm \frac{r_{j}}{2} s, \quad 0<s<1
$$

We believe our model represents a more accurate calculation by avoiding all of these assumptions. The first assumption is obviated by replacement of the parent with two daughter dipoles in all cases. This assumption becomes questionable when the parent splits into a daughter of roughly the same logarithmic size, for example when an $i=0$ parent splits into two $j=1$ daughters. In this case it is not accurate to maintain the parent since neither of the daughters are the same size. In fact, most allowed splittings are of this nature since a splitting where $j-i$ is large is unlikely to pass the $\kappa$ cutoff condition 152 . Possibly this difference accounts for our 1D wavefront velocity being higher than that of [24] (shown in figure 32), as sizes can be driven downwards faster when parents are removed and replaced by two smaller dipoles. Assumption 2 is not present in our model, since splittings like that shown in figure 16 left create dipoles that are not equal to $B^{-i}$ for any $i$. Assumption 3 reasonable in the collinear and infrared limits, but again, if $|j-i|$ is small then it is not accurate.

Assumption 4 is not necessary in our model because the splitting kernel determines the impact parameters of all daughter dipoles.
In summary, the splittings most relevant to driving the saturation front forward are those between similarly sized parent and daughter dipoles. Thus, it is important to handle these splittings accurately. We believe our model succeeds in this respect, and that it is therefore a more accurate model of dipole evolution than those previously wrought.

## 7 Final Summary

In this final chapter, we will more or less repeat was has already been said as concisely as possible. In chapter 2 we saw that a Regge trajectory with intercept greater than 1 called the Pomeron was needed to explain the rise of hadronic cross-sections. We then gave an account of how a pQCD calculation in the form of an infinite gluon ladder diagram could account for such a trajectory. In chapter 3 we introduced the dipole formulation for calculating cross-sections such as $\gamma^{*} p \rightarrow X$. In this picture, the virtual photon dissociates into a quark-antiquark pair which then interacts with the initial state hadron. Using this picture, Mueller showed that evolution of the target with increasing energy could be viewed as a highly occupied Fock state called an onium. Colorless dipoles comprise these states, which form due to soft gluon emissions. Using the wavefunction for the onium state, Mueller derived an integral equation which was equivalent to the BFKL equation found via the gluon ladder diagram, albeit the result of a much simpler calculation.

Although the BFKL equation correctly predicts dipole density growth in the dilute regime, in chapter 4 we explain that the eventual violation of unitarity with increasing $s$ necessitates a nonlinear growth taming term. This is provided by the BK equation, which adds a $-N^{2}$ term to the evolution equation, providing the desired effect. It was later shown by Munier and Peschanski that the BK equation belongs to the universality class of the FKPP equation, familiar from reaction-diffusion dynamics. This conceptual framework allowed the phenomenon of geometric scaling to be viewed as a traveling wave whose front is the logarithm of the saturation scale. This front moves with a group velocity equal to the minimum phase velocity of a wave packet in Mellin space, a condition that can be found by matching conditions in the dilute and saturation regions.

In Part II, we move on to describe a model based on the classical branching kernel of the BFKL equation and a saturation mechanism. Both the collinear and infrared parts of the kernel are taken into account. Saturation is checked by the program efficiently through the use of the red-black tree data structure. A full 2D implementation of the model as well as a 1D variant and a smooth interpolation between 1D and 2D are introduced. Data on wavefront asymptotics and correlations in impact parameter are presented and contrasted with an earlier work based on a 1D model. Finally, analytical calculations of the wavefront asymptotic velocity are compared with the
data.


Figure 33: Comparison of velocity and variance between $\rho_{\max }=20$ and $\rho_{\max }=50$.

## 8 Appendix

### 8.1 Dependence of the model on $\rho_{\max }$

During the final defense of this manuscript, the question was raised whether the length cutoff $r_{\text {min }}$, which in logarithmic coordinates is $\rho_{\max }:=\log _{B} \frac{1}{r_{\min }}$, in the divergent integral (134) has any effect on the results of the model. Analytically, we can see from (67) that the BFKL equation in Mellin space does not have a cutoff dependence. In fact, the lower size bound $\rho$ cancels in 62). Still, it may be asked whether this analytical cancellation applies to the model. I will demonstrate in several ways that the model does not have a strong dependence on $\rho_{\max }$ as long as it is sufficiently large.

### 8.1.1 Brute force model check

Running the model with different values of $\rho_{\max }$ is one way of checking for a possible dependence. For technical reasons explained before, it is not convenient to have $\rho_{\max } \gtrsim 50$, but we may check smaller values. Figure 33, for example, compares $\rho_{\max }=20$ and $\rho_{\max }=50$. Over 30 powers of the logarithmic base, the change in velocity and variance is small, although it appears the front velocity is slightly higher for the $\rho_{\max }=20$ case. This may be due to the change in relative splitting probabilities between near-size and far-size splittings. However, we believe that for sufficiently large $\rho_{\max }$ the artifact of higher front velocities disappears, as we now explain.

### 8.1.2 Lifetime dependence

Using the collinear branching probability approximation made in (169), we can write

$$
\begin{equation*}
\left.\frac{d P_{i \rightarrow j}}{d Y}\right|_{j>i} \approx 2\left(1+B^{-2(j-i)}\right) \tag{183}
\end{equation*}
$$

Making the fixed impact parameter approximation and multiplying by $\left(\frac{r_{j}}{r_{i}}\right)^{2}=$ $B^{-2(j-i)}$ as an estimate of the probability that the daughter $j$ will be created near the probe location,

$$
\left.\frac{d P_{i \rightarrow j}}{d Y}\right|_{j>i} \approx 2\left(B^{-2(j-i)}+B^{-4(j-i)}\right)
$$

Now finding the total probability of a size $i$ dipole splitting collinearly,

$$
\begin{equation*}
\frac{d P_{i}}{d Y} \approx \sum_{j=i+1}^{\rho_{\max }-1} 2\left(B^{-2(j-i)}+B^{-4(j-i)}\right) \tag{184}
\end{equation*}
$$

Observe this sum is convergent as $\rho_{\max } \rightarrow \infty$. Also, because it converges quickly, the effective dipole splitting rates (and lifetimes) are not highly sensitive to the exact value chosen for $\rho_{\max }$, as long as $\rho_{\max } \gg \rho_{s}(Y)$. By "effective", we mean the splittings that will affect the amplitudes measured at a particular impact parameter, which we estimated by adjusting the splitting probability by $B^{-2(j-i)}$.

### 8.1.3 Analytical check of BK equation using model constructs

We can explicitly check the BK equation (95) within the model construct to verify insensitivity to $\rho_{\max }$. To do so, we want to investigate the collinear part of the integral from limits 0 to $r_{\text {min }}$, which in logarithmic coordinates $\rho:=\log _{B} \frac{1}{x}$ become, respectively, $\infty$ and $\rho_{\max }$. Writing the BK equation using logarithmic coordinates at some impact parameter and using (183),

$$
\begin{equation*}
\partial_{Y} N_{i}(Y)=\sum_{j=\rho_{\max }}^{\infty} 2\left(1+B^{-2(j-i)}\right)\left[N_{f(i, j)}(Y)+N_{j}(Y)-N_{i}(Y)+N_{f(i, j)}(Y) N_{j}(Y)\right] \tag{185}
\end{equation*}
$$

where $i:=\log _{B} \frac{1}{x_{01}}, j:=\log _{B} \frac{1}{x_{12}}$, and

$$
\begin{aligned}
f(i, j) & :=\log _{B} \frac{1}{x_{02}} \\
& =\log _{B} \frac{1}{\sqrt{x_{01}^{2}+x_{12}^{2}-2 x_{01} x_{12} \cos \phi}} \\
& =\frac{1}{2} \log _{B} \frac{B^{2 i}}{1+B^{2(j-i)}-2 B^{(j-i)} \cos \phi} \\
& \approx \frac{1}{2} \log _{B}\left[B^{2 i}\left(1+B^{-2(j-i)}\right)\right]
\end{aligned}
$$

Assume that $Y$ is small enough such that $\rho_{\max } \gg \rho_{s}(Y)$. This is required for the validity of the model, as the wavefront must "fit" within the allotted logarithmic domain. Then $j \gg \rho_{s}(Y)$ and $N_{j}(Y) \approx 0$ far ahead of the saturation front. Also, because we are in the collinear region, $j \gg i$, assuming $\rho_{\max }$ is large enough that this is possible, and thus $f(i, j) \approx i$. Therefore, we see that with a sufficiently large $\rho_{\max }$, the term in brackets in 185 is approximately 0 . Further increasing $\rho_{\max }$ will have little effect on $\partial_{Y} N_{i}(Y)$.

### 8.2 2D Code

```
/*
2D Dipole Simulation
Author: Matt Haley
Versions:
2: uses red black tree removal
3: uses openmp
-
*/
#include <iostream>
#include < fstream>
#include <sstream>
#include<<cstdlib>
#include <ctime>
#include <cmath>
#include<vector>
#include <omp.h>
#include < stdio.h>
#include < stdlib.h>
using namespace std;
#include "datastructs/RedBlackTree4.h"
#include<codecogs/stats/dists/discrete/discrete/randomsample.h>
// Declare global variables
const double B=2;
const double delta=1;
const double pi=3.1415926535;
const double epsilon=pow(10.0, -14);
```

```
double r( const int & i )
{
    return pow(B, -i*delta);
}
bool areSame(double a, double b)
{
    return abs(a - b)<epsilon;
}
int main()
{
    // Seed random generator and make first call (predictable)
    srand ((unsigned) time (0));
    rand( );
    // Declare Input Vars -- all will be shared among threads and so should be const
    const double Y_max=3;
    const double Y1 =1,Y2=2,Y3=3; // output at these Y
    const double delta_ Y=.1;
    const int numEvents = 700;
    const double kappa_cutoff = pow(10.0, -1.0);
    //kappa_cutoff = 0; // disable cutoff
    const double b_probe=0;
    const double b_probe2=0; // make b_probeN=b_probe for faster runs at central IP
    const double b_probe3=0;
    const double b_probe4=0;
    const double b_probe5=0;
    //const double b_probe2=pow(10.0,-6.0); // make b_probeN=b_probe for faster runs at central IP
    //const double b_probe3=pow(10.0, -4.0);
    //const double b_probe4=pow(10.0, -2.0);
    //const double b_probe5=pow(10.0,-1.0);
    const int N_sat=25;
    const int N__initial=N_sat;
    //const int \overline{N_max=4*dōuble(N_sat)/pow(kappa_cutoff,2); // max number of dipoles of a given size}
    //cout << "N_max = " << N_max << endl;
    // double alphha_s=1;
    // N_sat=delta/alpha_s^2;
    const double probFactor = 2*log(B)/(2*pi);
    const int rho_min = 0;
    const int rho max = 40;
    //const int rho_max = 50;
    const int n_azimuth = 12;
    const double dphi = 2*pi/n_azimuth;
    // Declare Other Vars
    // using array for probability matrix instead of vector for multidimensionality
    // first entry of last dimension is sum over theta
    double prob_itoj[rho_max + 1][rho_max +1][n__azimuth+1] = {{{0}}};
    // initialize random generator, generate one value (predictable)
    RandGen gen;
    gen.RandInt(10);
    // determine discrete probability matrix for i>>j
    for( int i=0; i<=rho_max; i++)
    {
        for( int j=0; j<=rho_max; j++ )
        {
            double kthTerm;
            if( j<i ) {
            for( int k=3; k<=n_azimuth-3; k++) { // k limits depend on n_azimuth_-here 60<k<300 deg
                    kthTerm = probFactor*dphi/(1+pow(B, 2*(i-j)) - 2*pow(B,i-j)*c\overline{os}(k*dphi));
                    prob_itoj[i][j][0] += kthTerm; // k=0 is total angular prob i->j
                    prob_itoj[i][j][k+1]= kthTerm; // prob for the kth angular bin
            }
        }
            else if( j=i ) {
                for( int k=2; k<=n_azimuth-2; k++) { // k limits depend on n_azimuth_-here 60<k<300 deg
                    kthTerm = probFactor*dphi/(1+ pow(B, 2*(i-j)) - 2*pow(B,i-j)*cos(k*dphi));
```

```
                prob_itoj[i][j][0] += kthTerm; // k=0 is total angular prob i 
                prob_itoj[i][j][k+1] = kthTerm; // prob for the kth angular bin
            }
        }
        else { // j>i
            for( int k=0; k<=n_azimuth - 1; k++) {
            kthTerm = probFactor*dphi/(1+ pow(B, 2*(i-j)) -2*pow(B,i-j)*\operatorname{cos}(k*dphi));
            prob_itoj[i][j][0] += kthTerm; // k=0 is total angular prob i 
            prob_itoj[i][j][k+1]=kthTerm; // prob for the kth angular bin
            }
        }
    }
}
// print probability matrix
cout << "Probability Matrix:" << endl;
cout<< endl;
for( int i=0; i<=rho_max; i++)
{
    for( int j=0; j<=rho_max; j++ ) {
        cout << prob_itoj[i][j][0] << " ";
    }
    cout << endl;
}
cout << endl;
// determine lifetimes
vector<double> sum(rho_max+1);
vector<double> lifetime(rho_max+1); // upper limit on rho_a is rho_max-1??
for( int i=0; i<=rho_max; i++ ) {
    sum[i] = 0;
    for( int j=0; j<=rho_max; j++ ) {
        sum[i] += prob_itoj[i][j][0];
        }
        lifetime[i] = 1/sum[i];
}
// output lifetimes
cout << "Lifetimes:" << endl;
for( int rho_a=rho_min; rho_a<=rho_max; rho_a++) {
    cout << 1/lifetime[rho_a] << endl;
}
// print probability matrix, fixed i and j, print angular probabilities
/*
cout<< endl;
int i2 = 0;
int j2 = 0;
cout << prob_itoj[i2][j2][0] << ": ";
for( int k=0; k<=n_azimuth-1; k++) {
    cout << prob_ito\overline{j}[i2][j2][k+1]<<"";
}
cout << endl;
i2 = 0;
j2 = 1;
cout<< prob_itoj[i2][j2][0] << ": ";
for( int k=0; k<n_azimuth; k++) {
    cout << prob_itoj[i2][j2][k+1]<< " ";
}
cout << endl << endl;
*/
vector< Stats:: Dists:: Discrete:: Discrete::RandomSample<double>* > prob_itoj_gen(rho_max);
vector< Stats:: Dists:: Discrete:: Discrete::RandomSample<double>* > prob__k_azimuth_gen(2*(rho_max
        -1)+1);
double passToGenij[rho_max][rho_max] = {{0}};
double passToGenk[2*(rho_max - 1) - +1][n_azimuth] = {{0}};
for(int i=0; i<=rho_max-1; i++) {
    for(int j=0; j<=rho_max - 1; j++) {
        passToGenij[i][j] = prob_itoj[i][j][0];
    }
```

```
}
for(int i=0; i<=rho_max-1; i++) {
    prob_itoj_gen[i]=new Stats:: Dists:: Discrete:: Discrete:: RandomSample<double >(rho_max,
            passToGenij[i], true, time(0)/MERSENNEDIV);
    //prob_itoj_gen[i]=new Stats::Dists::Discrete::Discrete::RandomSample<double>(rho_max,
                prob_itoj[i], true, 0.3416);
}
for(int j=0; j<=2*(rho_max-1); j++) {
    if ( j<=rho_max-2 ) {
        for(int k=0; k<=n_azimuth - 1; k++) {
            passToGenk[j][k] = prob_itoj[rho_max - 1][j][k+1]; // last row of prob matrix
        }
    }
    else { // j>rho_max-2, where the rho_max-1 entry is for i>>i
        for(int k=0; \overline{k}<=n_azimuth - 1; k++) - {
            passToGenk[j][k] = prob_itoj[0][j][k+1]; // first row of prob matrix
        }
    }
}
for(int j=0; j<=2*(rho_max-1); j++) {
    prob_k_azimuth_gen[j]=new Stats:: Dists:: Discrete:: Discrete:: RandomSample<double>(n_azimuth
        passToGenk[j], true, time(0)/MERSENNEDIV);
}
cout << "Y_max = " << Y_max << ", kappa_cutoff = " << kappa_cutoff <<
        ", events = " << numEvents << endl;
cout << "probFactor = " << probFactor << endl;
// end serial code initializers
#pragma omp parallel // clear contents of output files
{
    int th_id = omp_get_thread_num();
    ofstream fileOutputStream, rho_sStream, TatProbeY1, TatProbeY2, TatProbeY3;
    stringstream ss;
    ss << th_id;
    string filename;
    filename = "rho_sCore" + ss.str() + ".dat";
    rho_sStream.open(filename.c_str()); // clears file contents
    ////rho_sStream << "numEvents=" << numEvents << endl;
    rho_sStream.close();
    filename = "TatProbeY1Core" + ss.str() + ".dat";
    TatProbeY1.open(filename.c_str()); // clears file contents
    ////TatProbeY1<< "numEvents= " << numEvents << endl;
    TatProbeY1.close();
    filename = "TatProbeY2Core" + ss.str() + ".dat";
    TatProbeY2.open(filename.c_str()); // clears file contents
    ////TatProbeY2 << "numEvents=" << numEvents << endl;
    TatProbeY2.close();
    filename = "TatProbeY3Core" + ss.str() + ".dat";
    TatProbeY3.open(filename.c_str()); // clears file contents
    ////TatProbeY3 << "numEvents=" << numEvents << endl;
    TatProbeY3.close();
}
// EVENT LOOP
cout << "Num procs = " << omp_get_num_procs() << endl;
#pragma omp parallel for
for( int event=1; event<=numEvents; event++ ) {
// initialize thread variables
double kappa,kappa2,kappa3,kappa4,kappa5;
double b01,b01x,b01y,x0x,x0y,x1x,x1y,x2x,x2y;
double x01x, x01y, x0 2x, x02y, x 12x, x12y;
double length_x01,x01hatx, x01haty,length_x02, length_x12;
double b02,b12,b02x,b02y,b12x,b12y;
```

```
double b02hatx, b02haty,b12hatx, b12haty, checkpointx, checkpointy;
int rho_x02, rho_x12;
double numsplits_i;
bool sizeRangex 02, sizeRangex 12, exceedkappax02, exceedkappax 12, unSatx02,unSatx02Lower,
        unSatx02Upper,
        unSatx12,unSatx12Lower,unSatx12Upper;
    double T[rho_max +1][5] = {{0}};
    double b;
    int rho_s,rho_sPrev;
    double angle, angle1, angle2;
    int count1, count2, count3, count4, count5, count6;
    count1= count 2= count 3=count4=\operatorname{count 5= count6=0;}
// stream vars
int th_id = omp_get_thread_num();
ofstream fileOutputStream,rho _sStream, TatProbeY1,TatProbeY2,TatProbeY3;
streambuf* sbuf = cout.rdbuf(); // make a copy of the cout stream buffer
stringstream ss;
ss << th_id;
string filename;
filename = "rho_sCore" + ss.str() + ".dat";
rho_sStream.open(filename.c_str(), ios::app); // appends to file contents
filename = "TatProbeY1Core" + ss.str() + ".dat";
TatProbeY1.open(filename.c_str(),ios::app); // appends to file contents
filename = "TatProbeY2Core" + ss.str() + ".dat";
TatProbeY2.open(filename.c_str(),ios::app); // appends to file contents
filename = "TatProbeY3Core" + ss.str() + ".dat";
TatProbeY3.open(filename.c_str(),ios::app); // appends to file contents
cout << "//////////////////// EVENT = " << event << ", core = " <<
        omp_get_thread_num() << " ////////////////////" << endl << endl;
// initialize n[i]
vector< RedBlackTree<double>* > n(rho_max+1);
for( int i=0; i<=rho_max; i++ ) {
    n[i]=new RedBlackTree<double >(-1000);
}
// populate the initial size dipoles
for( int k_b=1; k_b<=N_initial; k_b++ )
{
    angle1 = 2*pi*gen.RandReal();
    angle2 = 2*pi*gen.RandReal();
    b01 = r(0)/2*gen.RandReal();
    b01x = b01*cos(angle1);
    b01y = b01*sin(angle1);
    x0x = b01x + r(0)/2* cos(angle 2);
    x0y = b01y +r(0)/2*sin(angle2);
    x1x = b01x - r(0)/2* cos(angle2);
    x1y = b01y - r(0)/2*sin(angle 2);
    n[0]-> insert(b01, b01x, b01y, x0x, x0y, x1x, x1y);
}
rho_s=rho_sPrev=0;
// Rapidity Loop
for( double Y=0; Y<=Y_max+epsilon; Y=Y+delta_Y ) {
    cout << "////////////// Y = " << Y << endl < << endl; // output progress
    for( int i=0; i<=rho_max - 1; i++ ) {
        numsplits_i=1/lifetime[i]*delta_Y*n[i]->size();
        if( numsplits_i != 0 ) {
            //cout<< "\overline{i}="<< i<< ", numsplits_i=" << numsplits_i<< endl;
            //cout<< "lifetime=" << lifetime[i]}<<", size=" << \overline{n[i]->size()}<<<< endl
            //cout<< endl;
        }
        for( int l=1; l<=numsplits_i; l++ ) {
            if( n[i]->size() > 0 ) { // only split if dipoles exist
            // choose a random dipole from column i to split
            n[i]->randElement(b01, b01x, b01y, x0x, x0y, x1x, x1y);
            // choose size j to split into
```

```
int j=int( prob_itoj_gen[i] ->genReal() );
// choose azimuth k to split into
int k = int( prob_k_azimuth_gen[j-i+(rho_max - 1)] - >genReal() );
// calculate x2
angle = 2*pi*double(k)/double(n_azimuth);
x01x = x1x - x0x;
x01y = x1y - x0y;
length_x01 = pow(pow (x01x,2)+pow(x01y,2) ,.5); // Pythagorean thm
x01hatx = x01x/length_x01;
x01haty = x01y/length_x01;
x 2x = -r(j)*(cos(angle)*x01hatx - sin(angle)*x01haty); // just rotation piece
x 2y = -r(j)*(sin(angle)*x01hatx + cos(angle)*x01haty); // just rotation piece
// choose which side of x01 to split off of
if(gen.RandInt (0,1)==1) {// split off of x1
    x 2x = x 2x + x x m;
    x2y=x2y+x1y;
}
else { // split off of xO
    x 2x = -x 2x + x0x;
    x2y=-x2y+x0y;
}
// choose IP to split into
b02x = (x0x + x 2x)/2.0;
b02y=(x0y+x2y)/2.0;
b02 = pow(pow(b02x,2) +pow(b02y,2),.5);
b12x}=(\textrm{x}1\textrm{x}+\textrm{x}2\textrm{x})/2.0
b12y=(x1y+x2y)/2.0;
b12 = pow(pow(b12x,2) +pow(b12y,2),.5);
x02x = x 2x - x0x;
x02y = x 2y - x0y;
length_x02 = pow(pow(x02x,2) +pow(x02y,2),.5); // Pythagorean thm
x12x = x 2x - x1x;
x12y=x2y-x1y;
length_x12 = pow(pow(x12x,2) +pow(x12y,2),.5); // Pythagorean thm
// insert new dipoles, round new dipoles to nearest log_2
////rho_x02 = floor(log(1/length_x02)/log(2) + .5);
////rho_x12 = floor(log(1/length_x12)/log(2) + . 5);
rho_x02-}= int(log(1/length_x02)/\operatorname{log}(\textrm{B})+.5)
rho_x12 = int(log(1/length_x12)/log(B) + . 5);
/*
cout << "x0={"<< x0x <<","<<x0y<<"'}, x1={"<< x1x<<"","<< x1y<<
    "}, x2={"<< x2x<<"," << x2y<< "}"><< endl;
cout<<" b01={"<< b01x<<"," << b01y<<"'}, b02={"<< b02x <<
    "," << b02y<< "}, b12={" << b12x<< "," << b12y<< "}" << endl;
*/
/*
cout<< "endpoint x coordinates: " << x0x <<"," << x1x <<"," << x2x << endl;
cout<< "endpoint y coordinates: "<< x0y<<","<< x1y<<","<< x2y<< endl;
cout<< "IP x coordinates:"<< b01x<<","<<b02x<<","<< b12x<< endl;
cout<< "IP y coordinates:"<< b01y<< ","<< b02y<<"","<< b12y<< endl;
cout<< "rho_x01=" << i<<", rho_x02=" << rho_x02 <<", rho_x12=" << rho_x12 << endl;
cout<< end\overline{l}
*/
// check various conditions before adding daughters x02 and x12
unSatx02=unSatx02Lower=unSatx0 2 Upper=unSatx12=unSatx12Lower=unSatx12Upper=0;
sizeRangex02 = (rho_x02>= 0) && (rho_x02<= rho_max);
sizeRangex12 = (rho_x12>= 0) && (rho_x12<= rho_max);
kappa =r(rho_x02)/abs(b02-b_probe);
kappa2 = r(rho_x02)/abs(b02-\overline{b}_probe2);
kappa3 =r(rho_x02)/abs(b02-b_probe3);
kappa4 = r(rho_x02)/abs(b02-b_probe4);
kappa5 = r(rho_x02)/abs(b02-b_probe5);
exceedkappax02 = (kappa > kappa_cutoff) || (kappa2> kappa_cutoff) || (kappa3 >
    kappa_cutoff)
    || (kappa4 > kappa_cutoff) || (kappa5 > kappa_cutoff);
```

```
kappa = r(rho_x12)/abs(b12-b_probe);
kappa2 = r(rho_x12)/abs(b12-b_probe2);
kappa3 = r(rho_x12)/abs(b02-b_probe3);
kappa4 = r(rho_x12)/abs(b02-b_probe4);
kappa5 = r(rho_x12)/abs(b02-b_probe5);
```



```
    kappa_cutoff)
    || (kappa4 > kappa_cutoff) || (kappa5 > kappa_cutoff);
if( sizeRangex02 &&& sizeRangex12 && exceedkappax02 && exceedkappax12 ) {
    // assume already saturated if size> N_max
    //if( ((n[rho_x02]->size() >= N_max) // (n[rho_x12]->size() >= N_max)) ) continue;
    // if other tests passed, do time consuming saturation tests in nested form (nesting
        saves computation)
    unSatx02 = ( n[rho_x02]->between2D(b02-r(rho_x02)/2,b02+r(rho_x02)/2,r(rho_x02),b02x,
        b02y)< N_sat );
    if( unSatx02 ) { // lower boundary x02
        b02hatx = b02x/b02;
        b02haty = b02y/b02;
        checkpointx = b0 2x-r(rho x02)/2*b0 2hatx;
        checkpointy = b0 2y-r(rho_x02)/2*b02haty;
        if( b02-r(rho_x02)/2>= 人 ) {
            unSatx02Lower = n[rho_x02]-> between2D(b02-r(rho_x02),b02,r(rho_x02), checkpointx,
                checkpointy) < N_
    }
    else { // b02-r(rho_x02)/\mathscr{2}<0
            unSatx02Lower = n[rho_x02]-> between2D(0,abs(b02-r(rho_x02)),r(rho_x02), checkpointx,
                checkpointy) < N_sat;
    }
    if (unSatx02Lower ) { // upper boundary x02
            checkpointx = b0 2x+r(rho x02)/2*b02hatx;
            checkpointy = b02y+r(rho_x02)/2*b02haty;
            unSatx02Upper = n[rho_x02]->between2D(b02,b02+r(rho_x02),r(rho_x02), checkpointx,
                checkpointy) < N_ _sat;
            if(unSatx02Upper ) { ///at b12
                unSatx12 = n[rho_x12]-> between2D(b12-r(rho_x12)/2,b12+r(rho_x12)/2,r(rho_x12),b12x
                    , b12y) < N_-sat;
                if(unSatx12) \ // lower boundary x02
                    b12hatx = b12x/b12;
                    b12haty = b12y/b12;
                    checkpointx = b12x-r(rho_x12)/2*b12hatx;
                    checkpointy = b12y-r(rho_x12)/2*b12haty;
                    if( b12-r(rho_x12)/2>= 人 ) {
                    unSatx12Lower = n[rho_x12]-> between2D(b12-r(rho_x12),b12,r(rho_x12),
                    checkpointx, checkpointy) < N_sat;
                    }
                    else { // b12-r(rho_x12)/2<0
                            unSatx12Lower = n[rho_x12]-> between2D(0, abs(b12-r(rho_x12)),r(rho_x12),
                                    checkpointx, checkpointy) < N_sat;
                    }
                    if (unSatx12Lower ) { // upper boundary x12
                        checkpointx = b12x+r(rho_x12)/2*b12hatx;
                        checkpointy = b12y+r(rho_x12)/2*b12haty;
                        unSatx12Upper = n[rho_x12]->between2D(b12,b12+r(rho_x12),r(rho_x12),
                        checkpointx, checkpointy) < N_sat;
                    } // upper boundary x12
            } // lower boundary x12
            } // at b12
        } // upper boundary x02
    } // lower boundary x02
} // sizerange and kappa check
//else{
// cout<< "veto sizerange or kappa:"<< sizeRangex02<<<","<< sizeRangex12 << ","<<
// exceedkappax02 <<","<< exceedkappax12 << endl;
//}
if( unSatx02 && unSatx02Lower &&& unSatx02Upper &&& unSatx12 && unSatx 12Lower &&
    unSatx12Upper ) {
    if(floor(b01*10000) != floor(pow( pow(b01x,2) +pow(b01y,2),.5 )*10000) ) {
        cout << "*******ALERT*******: " << b01 << " " << pow( pow(b01x,2)+pow(b01y,2) ,.5 ) <<
            endl << endl << endl << endl;
```







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```









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``` 488 489
490 490
491 492 493 494 495 496 496
497 498 499
```

```
            }
```

            }
            ///n[i]->printTreeVector();
            ///n[i]->printTreeVector();
            n[i]->remove(b01); // remove parent
            n[i]->remove(b01); // remove parent
            n[rho_x02]->insert(b02,b02x,b02y,x0x, x0y, x2x, x2y); // add b02
            n[rho_x02]->insert(b02,b02x,b02y,x0x, x0y, x2x, x2y); // add b02
            n[rho_x12]->insert(b12,b12x,b12y,x1x, x1y, x2x,x2y); // add b12
            n[rho_x12]->insert(b12,b12x,b12y,x1x, x1y, x2x,x2y); // add b12
            // cout << "x0x=" << x0x<<", x1x=" << x1x<<", x2x=" << x2 x << endl;
            // cout << "x0x=" << x0x<<", x1x=" << x1x<<", x2x=" << x2 x << endl;
            //cout<<"x0y="<<x0y<<", x1y="<< x1y<<", x2y="<< x2y<< endl << endl;
            //cout<<"x0y="<<x0y<<", x1y="<< x1y<<", x2y="<< x2y<< endl << endl;
            //cout<< "entry rho_x02=" << rho_x02 <<":" << b02 <<"," << b02x <<"," << b02y <<
            //cout<< "entry rho_x02=" << rho_x02 <<":" << b02 <<"," << b02x <<"," << b02y <<
                "," << xOx << "," << xOy
                "," << xOx << "," << xOy
            // <<","<<x2x<<"," << x2y<< endl;
            // <<","<<x2x<<"," << x2y<< endl;
            //cout<< "entry rho_x12="<< rho_x12<<":"<< b12<<","<< b12x<<","<< b12y<<
            //cout<< "entry rho_x12="<< rho_x12<<":"<< b12<<","<< b12x<<","<< b12y<<
                "," << x1x<<"," << x1y
                "," << x1x<<"," << x1y
            // <<"," << x2x<<"," << x2y<< endl;
            // <<"," << x2x<<"," << x2y<< endl;
            //cout << "removal i=" << i <<"," " << b01<< "," << b01x <<""," << b01y << "," << x0x
            //cout << "removal i=" << i <<"," " << b01<< "," << b01x <<""," << b01y << "," << x0x
                <<"," << xOy
                <<"," << xOy
            // <<"," << x1x<<","<< x1y<< endl << endl;
            // <<"," << x1x<<","<< x1y<< endl << endl;
        } // saturation check
        } // saturation check
        //else{
        //else{
        // cout<<"veto saturation: "<<unSatx02 <<","<<unSatx02Lower <<"," << 
        // cout<<"veto saturation: "<<unSatx02 <<","<<unSatx02Lower <<"," << 
            unSatx02Upper<<
            unSatx02Upper<<
        // ", "<<unSatx12<<","<<unSatx12Lower<<","<<unSatx12Upper<<endl;
        // ", "<<unSatx12<<","<<unSatx12Lower<<","<<unSatx12Upper<<endl;
        //}
        //}
    } // tree size check
    } // tree size check
    } // dipole creation (numsplits)
    } // dipole creation (numsplits)
    } // i loop
} // i loop
// output amplitude
// output amplitude
for(int i=0; i<=rho_max; i++) {
for(int i=0; i<=rho_max; i++) {
for(int j=0; j<= 4; j++) {
for(int j=0; j<= 4; j++) {
if( j==0 ) b=0;
if( j==0 ) b=0;
else if( j==1) b=pow (10.0, -6);
else if( j==1) b=pow (10.0, -6);
else if( j==2) b=pow (10.0, -4);
else if( j==2) b=pow (10.0, -4);
else if( j==3 ) b=pow (10.0, -2);
else if( j==3 ) b=pow (10.0, -2);
else b=pow (10.0, -1); // j==4
else b=pow (10.0, -1); // j==4
T[i][j] = (double) n[i]->between2D(b-r(i)/2,b+r(i)/2,r(i),b,0)/( (double) N_sat);
T[i][j] = (double) n[i]->between2D(b-r(i)/2,b+r(i)/2,r(i),b,0)/( (double) N_sat);
}
}
}
}
if( areSame(Y,Y1) || areSame(Y,Y2) || areSame(Y,Y3) ) {
if( areSame(Y,Y1) || areSame(Y,Y2) || areSame(Y,Y3) ) {
cout << "OUTPUTTING AMPLITUDE" << endl << endl;
cout << "OUTPUTTING AMPLITUDE" << endl << endl;
if( areSame(Y,Y1) ) {
if( areSame(Y,Y1) ) {
for(int i=0; i<=rho_max; i++) {
for(int i=0; i<=rho_max; i++) {
TatProbeY1 << i << " " << T[i][0] << endl;
TatProbeY1 << i << " " << T[i][0] << endl;
}
}
TatProbeY1 << "end_of_event=" << event << endl;
TatProbeY1 << "end_of_event=" << event << endl;
}
}
if( areSame(Y,Y2) ) {
if( areSame(Y,Y2) ) {
for(int i=0; i<=rho_max; i++) {
for(int i=0; i<=rho_max; i++) {
TatProbeY2 << i << " " << T[i][0] << endl;
TatProbeY2 << i << " " << T[i][0] << endl;
}
}
TatProbeY2 << "end_of_event=" << event << endl;
TatProbeY2 << "end_of_event=" << event << endl;
}
}
if( areSame(Y,Y3) ) {
if( areSame(Y,Y3) ) {
for(int i=0; i<=rho_max; i++) {
for(int i=0; i<=rho_max; i++) {
TatProbeY3 << i << " " << T[i][0] << endl;
TatProbeY3 << i << " " << T[i][0] << endl;
}
}
TatProbeY3 << "end_of_event=" << event << endl;
TatProbeY3 << "end_of_event=" << event << endl;
}
}
/*for(int i=0; i<=rho_max-1; i++) {
/*for(int i=0; i<=rho_max-1; i++) {
fileOutputStream<<T[i]}<<\mathrm{ endl;
fileOutputStream<<T[i]}<<\mathrm{ endl;
}*/
}*/
///fileOutputStream.close();
///fileOutputStream.close();
//cout.rdbuf(sbuf); // reassign cout to console output
//cout.rdbuf(sbuf); // reassign cout to console output
} // end output
} // end output
// front position at central IP
// front position at central IP
rho_sStream << Y << " ";
rho_sStream << Y << " ";
for( int j=0; j< = 4; j++) {
for( int j=0; j< = 4; j++) {
for(int i=0; i<=rho_max; i++) {
for(int i=0; i<=rho_max; i++) {
if( T[i][j]>=.5 ) rho_s=i ;
if( T[i][j]>=.5 ) rho_s=i ;
}

```
    }
```

```
        rho_sStream << rho_s << " ";
    }
    rho_sStream << endl;
} // Y loop
fileOutputStream.open("TAmplitude.dat");
fileOutputStream << "b,b01x, b01y, x0x, x0y, x1x,x1y,kappa=" << kappa_cutoff << ",Y_max=" << Y_max
        <<
            ",numEvents=" << numEvents << ", probFactor=" << probFactor << endl;
cout.rdbuf(fileOutputStream.rdbuf()); // redirect cout to the output file stream
for( int i=0; i<=rho_max; i++ ) {
    ///cout<< "i="<< i<< endl;
    ///n[i]->printTreeVector();
}
cout.rdbuf(sbuf); // reassign cout to console output
fileOutputStream.close();
rho_sStream << "end_of_event=" << event << endl;
rho_sStream.close();
TatProbeY1.close();
TatProbeY2.close();
TatProbeY3.close();
} // EVENT LOOP, threads rejoin
// compile data
int numThreads = omp_get_num_procs();
ofstream fileOutputStream,rho_sStream,TatProbeY1,TatProbeY2,TatProbeY3;
ifstream input;
stringstream ss;
rho_sStream.open("rho_s.dat");
rho_sStream << "numEvents= " << numEvents << endl;
TatProbeY1.open("TatProbeY1.dat");
TatProbeY2.open("TatProbeY2.dat");
TatProbeY3.open("TatProbeY3.dat");
TatProbeY1 << "numEvents= " << numEvents << endl;
TatProbeY2 << "numEvents= " << numEvents << endl;
TatProbeY3 << "numEvents= " << numEvents << endl;
for( int i=0; i<numThreads; i++ ) {
    ss.str(""); // empty the string
    ss << i;
    string filename, data;
    filename = "rho_sCore" + ss.str() + ".dat";
    input.open(filename.c_str());
    if( !input.fail() ) {
        while( getline(input,data) ) {
            rho_sStream << data << endl;
        }
    }
    else cout << "Error: cannot open file " << filename << endl;
    input.close();
    filename = "TatProbeY1Core" + ss.str() + ".dat";
    input.open(filename.c_str());
    if( !input.fail() ) {
        while( getline(input,data) ) {
            TatProbeY1 << data << endl;
        }
    }
    else cout << "Error: cannot open file" << filename << endl;
    input.close();
    filename = "TatProbeY2Core" + ss.str() + ".dat";
    input.open(filename.c_str());
    if( !input.fail() ) {
        while( getline(input,data) ) {
            TatProbeY2 << data << endl;
        }
    }
    else cout << "Error: cannot open file" << filename << endl;
    input.close();
```

```
        filename = "TatProbeY3Core" + ss.str() + ".dat";
        input.open(filename.c_str());
        if( !input.fail() ) {
            while( getline(input, data) ) {
                TatProbeY3 << data << endl;
        }
    }
    else cout << "Error: cannot open file" << filename << endl;
        input.close();
    }
    rho_sStream.close();
    TatProbeY1.close();
    TatProbeY2.close();
    TatProbeY3.close();
    cout << "DONE, Y_max = " << Y_max << ", kappa_cutoff= " << kappa_cutoff <<
    ", events = " << numEvents << endl;
    cout << "probFactor = " << probFactor << endl;
    ///cout << "saturation veto counts:" << count1<<"," << count2 <<", " << count3 << ", " <<
    /// count4 <<"," << count5<<"," << count6<< endl;
    cout << flush;
    return 0;
}
```


### 8.3 2DR Code Snippet

```
// 2DR changes
x1xPrime = b12x + length_x m 2/2*x01hatx;
x 2xPrime = b 12x - length_x x 2/2*x01hatx;
x0xPrimePrime = b 02x - length_x02/2*x01hatx;
x 2xPrimePrime = b 0 2x + length_x x02/2*x01hatx;
b02y = 0;
b12y = 0;
b02 = abs(b02x);
b12 = abs(b12x);
```


### 8.4 2DSR Code Snippet

```
// 2DSR changes
s = gen.RandReal (0,1);
smallerRho = min(rho_x02,rho_x12);
stripwidth = stripFactor*r(smallerRho);
if(abs(x2y) > stripwidth ) {
    if( x 2y<0) s = - s; // project to the correct side of the x-axis
    x2xPrime = x 2x;
    x2yPrime = s*stripwidth;
    if( length_x02>abs(x0y-x2yPrime) ) {
        if( x0x > x2xPrime ) { // x0 slides up x-axis
            x0xPrime = x 2xPrime + pow(pow(length_x02,2)-pow(x0y-x2yPrime, 2),.5);
        }
            else { // x0x<= x2xPrime, xO slides down x-axis
            x0xPrime = x 2xPrime - pow(pow(length_x02, 2)-pow(x0y-x2yPrime, 2),.5);
        }
    }
    else x0xPrime = x0x;
    if( length_x12>abs(x1y-x2yPrime) ) {
        if( x1x > x2xPrime) { // x1 slides up x-axis
            x1xPrime = x 2xPrime + pow(pow(length_x12, 2)-pow(x1y-x2yPrime, 2),.5);
        }
        else { // x1x<= x2xPrime, x1 slides down x-axis
            x1xPrime = x 2xPrime - pow(pow(length_x12, 2)-pow(x1y-x2yPrime, 2),.5);
        }
    }
    else x1xPrime = x1x;
    if( x0xPrime!=x0xPrime || x1xPrime!=x1xPrime ) {
```

```
        cout << "********ALERT********" << endl;
    }
    x0yPrime = x0y;
    x1yPrime = x1y;
    // redefine impact parameters to primed ones
    b02x = (x0xPrime+x2xPrime)/2.0;
    b02y = (x0yPrime+x2y Prime)/2.0;
    b02 = pow(pow(b02x,2)+pow(b02y,2),.5);
    b12x = (x1xPrime+x2xPrime)/2.0;
    b12y = (x1yPrime+x2yPrime)/2.0;
    b12 = pow(pow(b12x,2)+pow(b12y,2),.5);
    // redefine coordinates to primed ones
    x0x = x0xPrime;
    x0y = x0yPrime;
    x1x = x1xPrime;
    x1y = x1yPrime;
    x2x = x2xPrime;
    x2y=x2yPrime;
}
```


### 8.5 RedBlackTree.h

```
    #ifndef RED_BLACK_TREE_H_
    #define RED_BLACK_TREE_H_
    #include "datastructs/dsexceptions.h"
    #include <iostream> // For NULL
#include <cstdlib>
#include "tapestry/randgen.h"
    // Red-black tree class
    //
    // CONSTRUCTION: with negative infinity object also
    // used to signal failed finds
    //
    // ********************PUBLIC OPERATIONS**********************
    // void insert( x ) --> Insert x
    // void remove( }x\mathrm{ ) --> Remove x (unimplemented)
    // Comparable find( x ) --> Return item that matches x
    // Comparable findMin( ) --> Return smallest item
    // Comparable findMax( ) --> Return largest item
    // boolean isEmpty( ) --> Return true if empty; else false
    // void makeEmpty( ) --> Remove all items
    // void printTree( ) --> Print tree in sorted order
// void printTree2( ) --> Print tree in tree order
// int size( ) --> Returns number of nodes in tree
// int between(x, y) --> Returns number of nodes with elements between x and y
// Comparable randElement(n) --> Returns a random tree element from the first n nodes
        // Node and forward declaration because g++ does
        // not understand nested classes.
    template <class Comparable>
    class RedBlackTree;
    template <class Comparable>
    class RedBlackNode
    {
            Comparable element;
            ///RedBlackNode *left;
            ///RedBlackNode *right;
            RedBlackNode *link[2]; // Left (0) and right (1) links
            int
                    red;
            double
            double
            by;
            double x0x;
            double x0y;
            double x1x;
```

    double \(x 1 y\);
    \(/ / c=O\) should be \(c=\) RedBlackTree \(<\) Comparable \(>:\) :BLACK
    // But Visual 5.0 does not comprehend it.
    RedBlackNode ( const Comparable \& theElement = Comparable( ) ,
                RedBlackNode \(* 1 \mathrm{t}=\) NULL, RedBlackNode \(* \mathrm{rt}=\) NULL,
                double thebx \(=\) double(), double theby \(=\) double (),
                double thex \(0 x=\) double(), double thex \(0 y=\) double(),
                double thexlx \(=\) double(), double thex1y \(=\) double(),
                int thered \(=1\) )
            : element (theElement ), red (thered), bx (thebx),
            by (theby), \(x 0 x(t h e x 0 x), x 0 y(t h e x 0 y), x 1 x(t h e x 1 x), \quad x 1 y(t h e x 1 y)\)
            \{
            link \([0]=1 \mathrm{t}\);
            link[1] \(=r t\);
            \(/ / /\) cout \(\ll\) "element=" \(\ll\) element \(\ll "\), mem of link[1]=" \(\ll \operatorname{link}[1] \ll e n d l\);
            \}
        friend class RedBlackTree \(<\) Comparable \(>\);
    \(\} ;\)
    template <class Comparable>
    class RedBlackTree
    \{
        public:
            explicit RedBlackTree( const Comparable \& negInf );
            RedBlackTree( const RedBlackTree \& rhs );
            ~RedBlackTree( );
            const Comparable \& findMin( ) const;
            const Comparable \& findMax ( ) const;
            const Comparable \& find ( const Comparable \& \(x\) ) const;
            bool isEmpty( ) const;
            void printTree( ) const;
    void printTree2( ) const;
void printTreeVector ( ) const;
int size( ) const;
int between (const Comparable \& lower, const Comparable \& upper) const;
int between2D (const Comparable \& lower, const Comparable \& upper,
const double \& ri, const double \& thebx, const double \& theby) const;
void randElement(const int $\mathcal{G} n$, Comparable $\mathcal{G}$ theElement,
double $\mathcal{G}$ thebx, double $\mathcal{G}$ theby,
double $\mathcal{G}$ thexOx, double $\mathcal{G}$ thexOy,
double $G$ thex $1 x$, double $G$ thex $1 y$ ) const;
void randElement ( Comparable \& theElement,
double $\&$ thebx, double $\&$ theby,
double \& thex0x, double \& thex $0 y$,
double \& thex1x, double \& thex1y) const;
void makeEmpty ( ) ;
//void insert (const Comparable $\varepsilon x$ );
int insert const Comparable \& $x$, const double \& $b x$,
const double \& by, const double \& x0x,
const double \& $x 0 y$, const double $\& x 1 x$,
const double \& $x 1 y$ );
int remove( const Comparable \& $x$ );
enum \{ BLACK, RED \};
const RedBlackTree \& operator $=($ const RedBlackTree \& rhs );
private:
RedBlackNode<Comparable $>$ *header; // The tree header (contains negInf)
const Comparable ITEM_NOT_FOUND;
RedBlackNode $<$ Comparable $>\bar{*}$ nullNode;
// Used in insert routine and its helpers (logically static)
RedBlackNode $<$ Comparable $>*$ current;
RedBlackNode $<$ Comparable $>*$ parent;
RedBlackNode $<$ Comparable $>*$ grand;
RedBlackNode $<$ Comparable $>$ *great ;

### 8.6 RedBlackTree.cpp

```
#include "RedBlackTree4.h"
#ifndef HEIGHT_LIMIT
#define HEIGHT_LIMIT 64/* Tallest allowable tree */
#endif
/**
    * Construct the tree.
    * negInf is a value less than or equal to all others.
    * It is also used as ITEM_NOT_FOUND.
    */
template <class Comparable>
RedBlackTree<Comparable>:: RedBlackTree( const Comparable & negInf )
    : ITEM_NOT_FOUND( negInf )
{
    nullNode = new RedBlackNode<Comparable >;
    nullNode }->\mathrm{ \link[0] = nullNode }->1\textrm{link}[1]=nullNode
    header = new RedBlackNode < Comparable>( negInf );
    header }->\mathrm{ link[0] = header }->\mathrm{ link [1] = nullNode;
    mySize = 0;
}
/**
    * Copy constructor.
    */
template <class Comparable>
RedBlackTree<Comparable >: RedBlackTree( const RedBlackTree<Comparable> & rhs )
    : ITEM_NOT_FOUND( rhs.ITEM_NOT_FOUND ), mySize(rhs.mySize)
{
    nullNode = new RedBlackNode<Comparable>;
    nullNode }->\mathrm{ link [0] = nullNode }->1\textrm{link}[1]=nullNode
    header = new RedBlackNode<Comparable>( ITEM_NOT_FOUND );
```

```
    header }->\mathrm{ link [0] = header }->\mathrm{ link [1] = nullNode;
    *this = rhs;
}
/**
    * Destroy the tree.
    */
template <class Comparable>
RedBlackTree<Comparable>::~ RedBlackTree( )
{
    makeEmpty( );
    delete nullNode;
    delete header;
}
/**
    * Remove item x from the tree
    * Not implemented in this version.
    */
template<class Comparable>
int RedBlackTree<Comparable >: 盾move( const Comparable & x )
{
    if (header }->\mathrm{ link[1] != nullNode ) {
        RedBlackNode<Comparable> head; /* False tree root */
        RedBlackNode<Comparable> *q, *p, *g; /* Helpers */
        RedBlackNode<Comparable> *f = nullNode; /* Found item */
            int dir = 1;
            /* Set up our helpers */
            q = &head;
            g}=\textrm{p}=\textrm{nullNode;
            q}->>link[0] = nullNode; // added so that looking above the root does not cause problem
            q}->>\operatorname{link[1] = header }->\mathrm{ link[1];
            /*
            Search and push a red node down
            to fix red violations as we go
            */
            while ( q->link[dir] != nullNode ) {
            int last = dir;
            /* Move the helpers down */
            g = p;
            p = q;
            q = q->link[dir];
            dir = q->element < x;
            /*
                Save the node with matching data and keep
                going; we'll do removal tasks at the end
            */
            if (q->element == x )
                f = q;
            /* Push the red node down with rotations and color flips */
            if (!is_red(q) && !is_red(q->>link[dir]) ) {
                if ( is_red ( q->lin\overline{k}[!dir]) )
                p = p-> link[last] = jsw_single ( q, dir );
            else if (!is_red (q->link[!dir]) ) {
                    RedBlackNode}<\mathrm{ Comparable }>*s=p>link[!last]
                    if (s != nullNode) {
                    if ( ! is_red(s }->\mathrm{ link[! last]) &&& !is_red(s s>link[last]) ) {
                        /* Color flip */
                        p}->\mathrm{ red = 0;
                        s }->\mathrm{ red = 1;
                        q}->\mathrm{ red = 1;
                    }
                    else {
```

```
                int dir2 = g-> link[1] == p;
                    if ( is_red (s->link[last]))
                    g->>link[dir2] = jsw_double ( p, last );
                    else if ( is_red ( s - link[!last] ) )
                    g->link[dir}2]=\mathrm{ jsw_single ( p, last );
                    /* Ensure correct coloring */
                q}->>\mathrm{ red = g }->\mathrm{ link[dir2] }>>\mathrm{ red = 1;
                g}->>\operatorname{link[dir2]-> link[0] }>>>\textrm{red}=0
                g}->\mathrm{ link[dir2] }>>\operatorname{link[1] }>>\mathrm{ red }=0
            }
            }
        }
        }
    } // end while
    /* Replace and remove the saved node */
    if ( f != nullNode) {
        ///tree->rel (f->element );
        f}->\mathrm{ -element = q }->\mathrm{ - element;
        f}->>bx=q->bx
        f}->>by=q->by
        f}->>0\textrm{x}=\textrm{q}->>\textrm{x}0\textrm{x}
        f}->>x0y=q->x0y
        f}->>x1x=q->x1x
        f}->>x1y=q->x1y
        p}->>\operatorname{link[p
            q}->>\operatorname{link[q}[>>1ink[0]== nullNode]
        delete(q);
        mySize--;
    }
    /* Update the root (it may be different) */
    header }->\mathrm{ link[1] = head.link[1];
    /* Make the root black for simplified logic */
    if (header }->\mathrm{ link[1] != nullNode )
        header }->\mathrm{ link[1] - >red = 0;
        ///--tree->size;
    }
    return 1;
}
/**
* Find the smallest item the tree
* Return the smallest item or ITEM_NOT_FOUND if empty.
*/
template <class Comparable>
const Comparable & RedBlackTree<Comparable>::findMin( ) const
{
    if( isEmpty( ) )
        return ITEM_NOT_FOUND;
    RedBlackNode<Comparable> *itr = header }->\mathrm{ link [1];
    while( itr }->\mathrm{ link[0] != nullNode )
        itr = itr }->\mathrm{ > link[0];
    return itr }->\mathrm{ - element;
}
/**
    * Find the largest item in the tree.
    * Return the largest item or ITEM_NOT_FOUND if empty.
    */
template<class Comparable>
const Comparable & RedBlackTree<Comparable>: findMax( ) const
```

```
{
    if( isEmpty( ) )
        return ITEM_NOT_FOUND;
    RedBlackNode<Comparable> *itr = header }->\mathrm{ >link[1];
    while( itr }->>>1ink[1] != nullNode )
        itr= itr }->\mathrm{ link[1];
    return itr }->\mathrm{ >element;
}
/**
    * Find item x in the tree.
    * Return the matching item or ITEM_NOT_FOUND if not found.
    */
template <class Comparable>
const Comparable & RedBlackTree<Comparable>::find( const Comparable & x ) const
{
    nullNode }->\mathrm{ -element = x;
    RedBlackNode<Comparable> *curr = header }->\mathrm{ \link [1];
    for( ; ; )
    {
        if( x < curr }->\mathrm{ - element )
            curr = curr }->\mathrm{ >link[0];
            else if( curr }->\mathrm{ >element < x )
                curr = curr->link[1];
            else if( curr != nullNode )
            return curr }->\mathrm{ - element;
            else
                return ITEM_NOT_FOUND;
    }
}
/**
    * Make the tree logically empty.
    */
template <class Comparable>
void RedBlackTree<Comparable > : makeEmpty( )
{
    reclaimMemory( header }->\mathrm{ link [1] );
    header }->\mathrm{ link[1] = nullNode;
}
/**
    * Test if the tree is logically empty.
    * Return true if empty, false otherwise.
    */
template <class Comparable>
bool RedBlackTree<Comparable>::isEmpty( ) const
{
    return header }->>1ink[1]= nullNode; 
}
/**
    * Print the tree contents in sorted order.
    */
template <class Comparable>
void RedBlackTree<Comparable >: : printTree( ) const
{
    if( header }->\mathrm{ link[1] == nullNode )
        cout<< "Empty tree" << endl;
    else
        printTree( header }->\operatorname{link[1] );
}
/**
    * Print the tree contents in binary tree order.
*/
```

```
template <class Comparable>
void RedBlackTree<Comparable>:: printTree2( ) const
{
    if(header }->\mathrm{ link[1] == nullNode )
        cout << "Empty tree" << endl;
    else
        printTree2( header }->\mathrm{ > link [1] );
}
/**
    * Print the tree contents in order sorted.
    */
template <class Comparable>
void RedBlackTree<Comparable > : printTreeVector( ) const
{
    if( header }->\mathrm{ link[1] == nullNode )
        cout << "Empty tree" << endl;
    else
        printTreeVector( header }->\mathrm{ >link [1] );
}
/**
    * Returns the number of nodes in the binary tree
*/
template <class Comparable>
int RedBlackTree<Comparable >: : size() const
{
    return mySize;
    //if(header>>link[1]== nullNode)
    // return 0;
    //else
    // return size(header }>>\operatorname{link[1] );
}
/**
    * Returns the number of node elements between lower and upper
    */
    template <class Comparable>
int RedBlackTree<Comparable >: between(const Comparable & lower, const Comparable & upper) const
{
    if( header }->\mathrm{ link[1] == nullNode)
        return 0;
    else
        return between(lower, upper, header }->\mathrm{ link [1]);
}
/**
* Returns the number of ode elements between lower and upper and within a radius r_i
*/
template <class Comparable>
int RedBlackTree<Comparable > : between2D(const Comparable & lower, const Comparable & upper,
            const double & ri, const double & thebx, const double & theby) const
{
    if( header }->\mathrm{ link[1] == nullNode)
        return 0;
    else
        return between2D(lower, upper, header }->\mathrm{ (link [1], ri, thebx, theby);
}
/**
    * Returns a random element between the 1st and nth nodes (in order)
    * R
template <class Comparable>
void RedBlackTree<Comparable>::randElement( Comparable & theElement,
            double & thebx, double & theby,
            double & thex0x, double & thex0y,
            double & thex1x, double & thex1y) const
{
    if( header }->\mathrm{ link[1] == nullNode) {
```

```
    cout << "error, randElement called on empty tree" << endl;
        return;
    }
    RandGen gen; // random number generator
    gen.RandInt( 1,mySize ); // first predictable
    int random_integer = gen.RandInt( 1,mySize );
    bool thedone = 0;
    return randElement( header }->\mathrm{ link[1], random_integer, theElement, thebx, theby, thex0x, thex0y,
        thex1x, thex1y, thedone );
}
/**
    * Deep copy.
    */
template <class Comparable>
const RedBlackTree<Comparable> &
RedBlackTree<Comparable > ::operator=( const RedBlackTree<Comparable> & rhs )
{
    if( this != &rhs )
    {
        makeEmpty( );
        header }->\mathrm{ link[1] = clone( rhs.header }->1ink[1] );
    }
    return *this;
}
/**
* Internal method to print a subtree t in sorted order.
    */
template <class Comparable>
void RedBlackTree<Comparable > :: printTree( RedBlackNode<Comparable> *t ) const
{
    if( t != t }->\mathrm{ link[0] )
    {
        printTree( t - link[0] );
        cout << t->element << endl;
        printTree( t }->\mathrm{ link[1] );
    }
}
/**
    * Internal method to print a subtree t in binary tree order.
*/
template <class Comparable>
void RedBlackTree<Comparable>:: printTree2( RedBlackNode<Comparable> *t ) const
{
    if( t != t }->\mathrm{ link[0] )
    {
        cout << t->element << endl;
        printTree2( t }->\mathrm{ link[0] );
        printTree2( t }->\mathrm{ link [1] );
    }
}
/**
    * Internal method to print a subtree t in sorted order.
    */
template <class Comparable>
void RedBlackTree<Comparable >: printTreeVector( RedBlackNode<Comparable> *t ) const
{
    if( t != t }->\mathrm{ link[0] )
    {
        printTreeVector( t }->\mathrm{ link [0] );
        cout << t->>element << " " << t ->bx << " " << t->>by << " " <<
            t->x0x << " " << t->x0y << " "<< t->>x1x << " " << t->x1y << endl;
        printTreeVector( t m link[1] );
    }
}
```

```
382
/*
    * Internal method to return the number of nodes in the binary tree
*/
template <class Comparable>
int RedBlackTree<Comparable > : : recursiveSize( RedBlackNode < Comparable> *t ) const
{
    if (t = = t }->\mathrm{ link[0])
        return 0;
    else
        return 1+ size( t }->>\operatorname{link[0] ) +\operatorname{size( t }->>\operatorname{link[1] );}
}
/**
    * Internal method to return the number of node elements between x and y
    */
template <class Comparable>
int RedBlackTree<Comparable > : between(const Comparable & lower, const Comparable & upper,
            RedBlackNode<Comparable> *t) const
{
    if( t = = t }->>\operatorname{link[0] )
        return 0;
    else if( t->element > lower && t->element < upper)
        return 1+between( lower, upper, t }->\mathrm{ link [0] ) +between( lower, upper, t }->\mathrm{ link [1]);
    else if( t->element > lower )
        return between( lower, upper, t }->\mathrm{ link [0]);
    else if( t->element < upper )
        return between( lower, upper,t }->>>link[1])
    else
    {
        cout << "error" << endl;
        return 0;
    }
}
/**
    * Internal method to return the number of node elements between x and y and within radius r_i
    */
    template <class Comparable>
int RedBlackTree<Comparable >: between2D (const Comparable & lower, const Comparable & upper,
    RedBlackNode<Comparable> *t, const double & ri, const double & thebx, const double & theby)
                const
{
    if( t == t }->>\operatorname{link[0] )
        return 0;
    else if( t }->\mathrm{ <element >= lower &&& t }->>\mathrm{ element <= upper) {
        ///if(t->hasSplit== 0 ) {
        double dist = pow(pow(thebx - t }->>\textrm{bx},2)+\mathrm{ pow(theby - t }->>by,2),.5)
        if( dist <= ri/2 ) { // check vector distance
            return 1+between2D(lower, upper, t }->\mathrm{ link [0], ri, thebx, theby) +
                between2D(lower, upper, t }->>\operatorname{link[1], ri, thebx, theby);
            }
            else { // not within vector distance, keep looking
            return 0+between2D(lower, upper, t }->1\textrm{link}[0],ri, thebx, theby)
                between2D(lower, upper, t }->\mathrm{ link[1], ri, thebx, theby);
            }
    }
        ///else {// already split, don't count for saturation
        /// return 0+between2D(lower, upper,t->link[0], ri, thebx, theby)+
        /// between2D(lower,upper, t->>ink[1], ri, thebx,theby);
        ///}
    else if( t }->\mathrm{ <element > lower )
            return between2D(lower, upper, t }->\mathrm{ link [0], ri, thebx, theby);
    else if( t->element < upper )
            return between2D(lower, upper, t }->\mathrm{ link [1], ri, thebx, theby);
    else
    {
        cout << "error, between2D failed" << endl;
            return 0;
    }
```

```
}
/**
    * Internal method to return the randomly chosen dipole
*/
template <class Comparable>
void RedBlackTree<Comparable>::randElement( RedBlackNode<Comparable> *t, int & countdown,
        Comparable & theElement,
        double & thebx, double & theby,
        double & thex0x, double & thex0y,
        double & thex1x, double & thex1y, bool & thedone) const
{
    if( thedone== 1) return;
    countdown--;
    if( t == nullNode ) {
        countdown++;
        return;
    }
    else if( countdown=0) {
        theElement = t }->\mathrm{ - element;
        thebx = t }->\mathrm{ bx;
        theby = t }->\mathrm{ by;
        thex0x = t }->\textrm{x}0\textrm{x}
        thex0y = t }->\textrm{x}0\textrm{y}
        thex1x = t }->\textrm{x}1\textrm{x}
        thex1y = t }->>x1y
        ///countdown = -1000;
        thedone = 1;
        return;
    }
    else {
            randElement( t > link[0], countdown, theElement, thebx, theby, thex 0x, thex0y, thex 1x, thex 1y,
                thedone );
            randElement( t > link[1], countdown, theElement, thebx, theby, thex0x, thex0y, thex 1x, thex1y,
                thedone );
            return;
    }
}
/**
    * Internal method to clone subtree.
    */
    template <class Comparable>
    RedBlackNode<Comparable> *
    RedBlackTree<Comparable > : clone( RedBlackNode<Comparable> * t ) const
{
    if( t == t }->\mathrm{ link[0] ) // Cannot test against nullNode!!!
        return nullNode;
    else
            return new RedBlackNode<Comparable>( t }->\mathrm{ <element, clone( t }->>>1ink[0] )
                clone( t }>>link[1] ), t->>color, t->>bx, t->>by
                t->>x0x, t->>x0y, t->>x1x, t->>x1y);
    }
    /**
    <summary>
    Performs a single red black rotation in the specified direction
    This function assumes that all nodes are valid for a rotation
    <summary>
    <param name="root">The original root to rotate around</param>
    <param name="dir">The direction to rotate (0=left, 1 = right)</param>
    <returns>The new root ater rotation</returns>
    <remarks>For jsw_rbtree.c internal use only</remarks>
*/
template <class Comparable>
RedBlackNode<Comparable> * RedBlackTree<Comparable > : j jsw_single( RedBlackNode<Comparable> *root ,
        int dir ) const
    {
    RedBlackNode<Comparable> *save = root }>>\mathrm{ link[!dir];
```

```
    root }->\mathrm{ link [! dir] = save }->\mathrm{ >link[dir];
    save}->\mathrm{ link[dir] = root;
    root }->\mathrm{ red = 1;
    save }->\mathrm{ red = 0;
    return save;
}
/**
    <summary>
    Performs a double red black rotation in the specified direction
    This function assumes that all nodes are valid for a rotation
    <summary>
    <param name="root">The original root to rotate around</param>
    <param name="dir">The direction to rotate (0 = left, 1 = right)</param>
    <returns>The new root after rotation</returns>
    <remarks>For jsw_rbtree.c internal use only</remarks>
*/
template <class Comparable>
RedBlackNode<Comparable> * RedBlackTree<Comparable > : jsw_double( RedBlackNode<Comparable> *root ,
        int dir ) const
{
    root }->\mathrm{ >link [!dir] = jsw_single ( root }->\mathrm{ link [!dir], !dir );
    return jsw_single ( root, dir );
}
/**
    <summary>
    Insert a copy of the user-specified
    data into a red black tree
    <summary>
    <param name="tree">The tree to insert into</param>
    <param name="data"> The data value to insert</param>
    <returns>
    1 if the value was inserted successfully,
    O if the insertion failed for any reason
    </returns>
*/
template<class Comparable>
int RedBlackTree<Comparable>::insert( const Comparable & x, const double & bx,
        const double & by, const double & x0x, const double & x0y, const double & x mx,
        const double & x1y )
{
    if( header }->\mathrm{ link[1] == nullNode) {
        /*
            We have an empty tree; attach the
            new node directly to the root
        */
        header }->\mathrm{ link[1] = new RedBlackNode<Comparable>( x, nullNode, nullNode, bx, by, x0x, x0y, x1x,
                    x1y );
            if (header }->\mathrm{ link[1] = nullNode ) {
            return 0;
        }
            else mySize++;
    }
    else {
        ///jsw_rbnode_t head = {0}; /* False tree root */
        //RedBlackNode<Comparable> head = new RedBlackNode<Comparable>;
        RedBlackNode<Comparable> head; /* False tree root */
        //RedBlackNode<Comparable> *head; /* False tree root */
        RedBlackNode<Comparable> *g, *t; /* Grandparent Eg parent */
        RedBlackNode<Comparable> *p, *q; //* Iterator & parent */
        int dir = 0, last = 0;
            /* Set up our helpers */
            t = &head;
```

```
    ///cout << "hi5" << endl;
    g = p = nullNode;
    q}=\textrm{t}->>\operatorname{link[1] = header }->\mathrm{ >link [1];
    ///cout<< "hi6"<< endl;
    /* Search down the tree for a place to insert */
    for ( ; ; ) {
        if ( q == nullNode) {
        /* Insert a new node at the first null link */
        p}->\mathrm{ >link[dir] = q = new RedBlackNode<Comparable>( x, nullNode, nullNode, bx, by, x0x, x0y,
                    x1x, x1y );
        if ( q == nullNode )
            return 0;
        else mySize++;
        }
        else if ( is_red ( q->link[0] ) && is_red ( q->link[1] ) ) {
        /* Simple red violation: color flip */
        q}->\mathrm{ red = 1;
        q}->>\operatorname{link[0] - > red = 0;
        q}->>\operatorname{link[1] -> red = 0;
        }
        if ( is_red ( q ) && is_red ( p ) ) {
        /* Hard red violation: rotations necessary */
        int dir2 = t }->\mathrm{ >link[1] == g;
        if ( q == p->link[last] )
            t->>link[dir2] = jsw_single ( g, !last );
        else
            t->>link[dir2] = jsw_double ( g, !last );
        }
        /*
        Stop working if we inserted a node. This
        check also disallows duplicates in the tree
        */
        if (q->element == x )
        break;
        last = dir;
        dir = q->element < x;
        /* Move the helpers down */
        if (g != nullNode)
        t = g;
        g = p, p = q;
        q = q->link[dir];
        }
        /* Update the root (it may be different) */
        header }->\mathrm{ link [1] = head.link [1];
    }
    /* Make the root black for simplified logic */
    header }->\mathrm{ link[1] -> red = 0;
    return 1;
}
///**
// * Insert item x into the tree. Does nothing if x already present.
// */
//template<class Comparable>
//void RedBlackTree<Comparable>::insert( const Comparable G x, const double G bx,
// const double छGby, const double छG xOx, const double छgx0y, const double G x fx,
// const double & x1y)
//{
// current = parent = grand= header;
```

```
// nullNode->>element = x;
//
while( current->element != x )
{
    great = grand; grand = parent; parent = current;
    current = x<current >element? current->>eft:current->>right;
        // Check if two red children; fix if so
    if(current->left }->\mathrm{ color == RED GGG current }->>\mathrm{ right }->\mathrm{ color = = RED )
            handleReorient( }x\mathrm{ );
}
    // Insertion fails if already present
if(current != nullNode)
    return;
current = new RedBlackNode<Comparable>( x, nullNode, nullNode, bx, by, x0x, x0y, x1x, x1y );
    // Attach to parent
if( }x<\mathrm{ parent->element)
    parent->>left = current;
else
    parent->right = current;
handleReorient( }x\mathrm{ );
//}
///**
// * Internal routine that is called during an insertion
// * if a node has two red children. Performs flip
// * and rotatons.
// * item is the item being inserted.
// */
//template <class Comparable>
//void RedBlackTree<Comparable >::handleReorient( const Comparable GG item)
//{
// // Do the color flip
// current->color= RED;
// current }>\mathrm{ left }->\mathrm{ color = BLACK;
// current }>\mathrm{ >right }->\mathrm{ color = BLACK;
//
// if( parent }->\mathrm{ color == RED ) // Have to rotate
// {
// grand->color= RED;
// if( item < grand->element != item < parent->element )
            parent = rotate( item, grand); // Start dbl rotate
    current = rotate(item, great );
    current->color=BLACK;
}
header }->\mathrm{ >right }->\mathrm{ color = BLACK; // Make root black
//}
//
///**
/ Internal routine that performs a single or double rotation.
// * Because the result is attached to the parent, there are four cases.
// * Called by handleReorient.
// * item is the item in handleReorient.
// * parent is the parent of the root of the rotated subtree.
// * Return the root of the rotated subtree.
// */
//template <class Comparable>
//RedBlackNode<Comparable> *
//RedBlackTree<Comparable>::rotate( const Comparable EG item,
// RedBlackNode<Comparable> *theParent ) const
//{
// if( item<theParent->element )
// {
// item<theParent->left >element?
            rotateWithLeftChild( theParent->left) : // LL
            rotate WithRightChild ( theParent }>>left) ; // L
            return theParent->left;
```

```
// }
// else
// {
// item<theParent->right->element?
// rotateWithLeftChild( theParent->right ) : // RL
// rotateWithLeftChild( theParent->right ) : // RL
// return theParent->right;
// }
//}
//
///**
// * Rotate binary tree node with left child.
// */
//template <class Comparable>
//void RedBlackTree<Comparable>::
//rotate WithLeftChild( RedBlackNode<Comparable> * G k2 ) const
//{
// RedBlackNode<Comparable> *k1 = k2->left;
// k2->left = k1->right;
// k1->right = k2;
// k2 = k1;
//}
//
///**
// * Rotate binary tree node with right child.
// */
//template <class Comparable>
//void RedBlackTree<Comparable >::
//rotateWithRightChild( RedBlackNode<Comparable> * छ k1 ) const
//{
// RedBlackNode<Comparable> *k2 = k1-> right;
// k1->right = k2->left;
// k2->left = k1;
// k1 = k2;
//}
/**
    * Internal method to reclaim internal nodes
    * in subtree t.
    */
template <class Comparable>
void RedBlackTree<Comparable >: reclaimMemory( RedBlackNode<Comparable> *t ) const
{
    if( t != t }->\mathrm{ link[0] )
    {
        reclaimMemory( t }->>>1ink[0] )
        reclaimMemory( t->link[1] );
        delete t;
    }
}
/**
    <summary>
    Checks the color of a red black node
    <summary>
    <param name="root">The node to check</param>
    <returns>1 for a red node, 0 for a black node</returns>
    <remarks>For jsw_rbtree.c internal use only</remarks>
*/
template <class Comparable>
int RedBlackTree<Comparable >: is_red ( RedBlackNode<Comparable> *root ) const
{
    return root != nullNode && root }->\mathrm{ red = 1;
}
```


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[^0]:    ${ }^{1}$ Fits to the data actually indicate the presence of two kinds of Pomeron: a "soft" Pomeron with behavior $s^{0.08}$ and a "hard" pomeron with behavior $s^{0.4}$. Because the soft Pomeron lies outside the reach of perturbative methods, we will only focus on the hard Pomeron. [5]

[^1]:    ${ }^{2}$ This is related to T'Hooft's observation that for $\mathrm{SU}(\mathrm{N})$, as $\mathrm{N} \rightarrow \infty$ planar graphs dominate over those of differing topology [49].

[^2]:    ${ }^{3}$ Technically, this would be the inverse Laplace transform of $T_{\omega}\left(Q x_{10}\right)$, but these transforms are related since $\left\{\mathcal{M}^{-1} T_{\omega}\right\}\left(e^{-Y}\right)=\frac{1}{2 \pi i} \int_{c} e^{\omega Y} T_{\omega} d \omega=\left\{\mathcal{L}^{-1} T_{\omega}\right\}(Y)$.

[^3]:    ${ }^{4}$ assuming degenerate trajectories for even and odd C-parity
    ${ }^{5}$ Interestingly, because the NLO correction is so substantial, Donnachie et. al. claim the perturbative ladder diagram calculation of the BFKL pomeron is suspect and that the correct value for the hard pomeron intercept provided by this calculation is probably a coincidence. See section 7.3 of [7] for details.

[^4]:    ${ }^{6}$ This is basically a rewriting of $\sqrt{22}$

[^5]:    ${ }^{7}$ N.B. We have made a trivial change to comply with more modern notation, $2 \chi_{\text {Mueller }}(\lambda=$ $2(1-\gamma))=\chi_{B F K L}\left(\gamma=1-\frac{\lambda}{2}\right)=: \chi(\gamma)=2 \psi(1)-\psi(1-\gamma)-\psi(\gamma) 44$. Thus the poles displayed in figure 12 are transformed like so: $\lambda=0 \rightarrow \gamma=1$ and $\lambda=2 \rightarrow \gamma=0$. Also, the saddle point at $\lambda_{s}=1 \rightarrow \gamma_{s}=\frac{1}{2}$. For the remainder of this manuscript, we mean " $\chi_{B F K L}$ " when we write " $\chi$ ".

[^6]:    ${ }^{8}$ Remember that $t \equiv Y$.

[^7]:    ${ }^{9}$ Note that " $Y$ " in this model is actually rapidity scaled by $\bar{\alpha}$. I.e $\bar{\alpha} Y \rightarrow Y$ throughout Part II.

[^8]:    ${ }^{10} \rho_{\text {max }}=50$ is chosen due to the fact that 64 -bit double precision binary floating-point numbers carry 1 bit of sign, 11 bits of exponent width, and 52 bits of significand precision. Thus, the maximum rounding error between two numbers, or machine epsilon, is $2^{-53}$. $\rho_{\max }$ should be kept well below 53.

[^9]:    ${ }^{11}$ In practice, to avoid creating $\rho_{\max }^{2}$ discrete probability distributions for azimuth selection, we note that 141) depends on $j-i$, which is bounded between $-\rho_{\max } \leq j-i \leq \rho_{\max }$. Thus we only need to create $2 \rho_{\max }+1$ discrete probability distributions.

[^10]:    ${ }^{12}$ Recall from the logarithmic size definition under 136 that $r_{i}=B^{-i}$.

[^11]:    ${ }^{13}$ The AVL tree is also sometimes used.

[^12]:    ${ }^{14}$ The terminus of a path.

[^13]:    ${ }^{15}$ unless $x_{02}^{2}<\left(x_{0 y}-x_{2 y}^{\prime}\right)^{2}$ or $x_{12}^{2}<\left(x_{1 y}-x_{2 y}^{\prime}\right)^{2}$.

[^14]:    ${ }^{16}$ The velocity values were averaged over $Y=1$ to $Y=3$.

[^15]:    ${ }^{17}$ The velocity values were averaged over $Y=1.5$ to $Y=3$.

