Galactic cosmic-ray propagation in the light of AMS-02: I. Analysis of protons, helium, and antiprotons

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We present novel constraints on cosmic-ray propagation in the Galaxy using the recent precise measurements of proton and helium spectra from AMS-02, together with preliminary AMS-02 data on the antiproton over proton ratio. To explore efficiently the large (up to eleven-dimensional) parameter space we employ the nested-sampling algorithm as implemented in the MultiNest package, interfaced with the Galprop code to compute the model-predicted spectra. We use VOYAGER proton and helium data, sampling the local interstellar spectra, to constrain the solar modulation potential. We find that the turbulence of the Galactic magnetic field is well constrained, i.e., \( \delta = 0.30^{+0.03}_{-0.02} \text{(stat)}^{+0.10}_{-0.04} \text{(sys)} \), with uncertainties dominated by systematic effects. Systematic uncertainties are determined checking the robustness of the results to the minimum rigidity cut used to fit the data (from 1GV to 5GV), to the propagation scenario (convection vs no convection), and to the uncertainties in the knowledge of the antiproton production cross section. Convection and reacceleration are found to be degenerate and not well constrained singularly when using data above 5GV. Using data above 1GV reacceleration is required, \( v_A = 25 \pm 2 \text{km/s} \), although this value might be significantly affected by the low-energy systematic uncertainty in the solar modulation. In a forthcoming companion paper, we investigate the constraints imposed by AMS-02 measurements on lithium, boron, and carbon.

INTRODUCTION

Cosmic-ray (CR) physics is on the verge of transition to a precision era thanks to the recently available data from PAMELA first, and more recently from the AMS-02 experiment on board the International Space Station. Thanks to these precise data, cracks in the standard minimal scenario start to appear. For example, a significant difference in the slopes of proton and helium, of about \( \sim 0.1 \) [1–3]), has been observed, while, from the standard CR acceleration scenario, no differences would be expected, at least for energies above 20-30 GeVs. The same measurements also find a break in the proton and helium rigidity spectra at about 300 GV. In this case, the feature can be accommodated with an extension of the standard scenario, and various explanations have been proposed [4–8].

Nonetheless, besides the above CR ‘anomalies’, the standard diffusion-reacceleration-convection scenario is, in the first place, not yet very well constrained. For example, estimates of the degree of turbulence in the Galactic magnetic field (encoded in the parameter conventionally indicated as \( \delta \)) range from the standard Kolmogorov turbulence (\( \delta = 0.33 \), [9, 10]) to Kraichnan (\( \delta = 0.5 \)) or plain diffusion (\( \delta = 0.6 \), [11]), up to \( \delta = 0.9 \) [12, 13]. Again, the new precise data offer the possibility to finally pin down the uncertainties in the parameters of the model.

In the following, we will thus analyze recently published proton [2], and helium [3] AMS-02 data together with preliminary AMS-02 data on the antiproton over proton ratio [14]. The analysis of the heavier nuclei, lithium, boron and carbon and comparison with the results from the analysis of this work is presented in a companion forthcoming article.

We also treat in a novel way the effect of solar modulation. Although we still use the force-field approximation, we do not assume any prior on the solar modulation potential, but we, instead, use recent VOYAGER data [15], sampling the interstellar unmodulated CR spectrum, to constrain the amount of solar modulation. The effect of solar modulation will be also studied applying different cuts on the minimum rigidity of the data used in the fit. Finally, we will also investigate the effect of uncertainties in the antiproton production cross section, following the recent redetermination from [16].

There is another well-known anomaly in CRs, namely the rising positron fraction observed by both PAMELA [17] and AMS-02 [18]. The rising is incompatible with the usual interpretation of positrons as secondaries produced by protons during their propagation. Although some attempt has been made to reconcile the positron fraction with the interpretation as secondaries through some modification of the propagation model [19], the generally accepted explanation requires a primary source of positrons, like pulsars or a nearby supernova remnant, or possibly, dark

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matter annihilation. In our study, we will assume the standard propagation scenario described above, and, as such, we would require a primary positron source to explain the observations. Nonetheless, even after including a primary source, it might be nontrivial to explain the positron fraction as well as the $e^+ + e^-$ spectrum, since the propagation of leptons is significantly affected by energy losses in the local radiation and magnetic field, while the local turbulence properties of the magnetic field can be different from the large scale average probed by nuclei. Although it would be, thus, interesting to cross-check the results of the study of propagation of nuclei with lepton spectra observations, ultimately, the above issues would make the comparison complicated and difficult to interpret. We will thus avoid these comparisons in the following and focus only on nuclei.

The work is structured as follows: The theoretical framework is discussed in SECTION I. The fit methodology is discussed in SECTION II. The results are presented in SECTION III, while we conclude in SECTION IV.

I. THEORY

The propagation of CR can be described by the well-known diffusion equation [20] for the particle density $\psi_i$ of species $i$ per volume and absolute value of momentum $p$

$$\frac{\partial \psi_i(x, p, t)}{\partial t} = q_i(x, p) + \nabla \cdot (D_{xx} \nabla \psi_i - V_i \psi_i)$$

$$+ \frac{\partial}{\partial p} p^2 D_{pp} \frac{\partial}{\partial p} \frac{1}{p^2} \psi_i - \frac{\partial}{\partial p} \left( \frac{dp}{dt} \psi_i - \frac{p}{3} (\nabla \cdot V) \psi_i \right) - \frac{1}{\tau_{f,i}} \psi_i - \frac{1}{\tau_{r,i}} \psi_i. \quad (1)$$

The various terms describe (i) spatial diffusion, usually assumed to be homogeneous and isotropic and thus described by the momentum-dependent diffusion coefficient $D_{xx}(p)$, (ii) convective winds, described by their velocity $V(x)$, (iii) diffusive reacceleration, parametrized as a diffusion in momentum space with coefficient $D_{pp}(p)$, (iv) continuous energy losses through the coefficient $dp/dt = \sum_k dp_k/dt$ which sums over all the various processes, $dp_k/dt$, through which the particles lose energy, (v) adiabatic energy losses, present if $V(x)$ has a nonzero divergence, and finally, catastrophic losses by (vi) decay or (vii) fragmentation, with decay and interaction times $\tau_r$ and $\tau_f$, respectively. The equation is typically solved assuming a steady state regime, meaning that $\psi_i$ does not depend on time and so the term on left-hand side is zero.

Diffusion is naturally expected to be an energy dependent process, with particles being less deflected by the magnetic fields with increasing energy, and thus diffusing faster. This process is usually modeled by a power law in rigidity $R = p/|Z|$ ([21]):

$$D_{xx} = \beta D_0 \left( \frac{R}{4 \text{GV}} \right)^\delta,$$  \quad (2)

where $\delta$ is the index of the power-law, $D_0$ the overall normalization, and $\beta = v/c$ the velocity of the CRs; we set the normalization scale at 4 GV. The constant for diffusive reacceleration $D_{pp}$ is usually related to the spatial diffusion $D_{xx}$ and to the velocity $v_A$ of Alfvén magnetic waves [22, 23] as

$$D_{pp} = \frac{4 (p v_A)^2}{3(2 - \delta)(2 + \delta)(4 - \delta) \delta D_{xx}}. \quad (3)$$

The amount of reacceleration is thus described in terms of the parameter $v_A$. Finally, convective winds are assumed to be constant and orthogonal to the Galactic plane $V(x) = \text{sign}(z) v_{0,c}$. We note that, in principle, this parametrization implies an unphysical discontinuity at $z = 0$. A smooth transition in the thin halo containing the sources (with size $\sim$0.2 kpc) would be more realistic. Nonetheless, since this parametrization has been widely employed in past works, we use it for the sake of comparison.

The source term $q_i(x, p)$ of primary CR is assumed to factorize into a species dependent normalization $q_{0,i}$, a space-depend part $q_{r,z}$ (where $r = \sqrt{x^2 + y^2}$ and $z$ are Galactocentric cylindrical coordinates), and a rigidity dependent part $q_R$:

$$q_i(x, p) = q_i(r, z, R) = q_{0,i} q_{r,z}(r, z) q_R(R). \quad (4)$$

We model the rigidity dependence as double broken power law with smooth transitions

$$q_R(R) = \left( \frac{R}{R_0} \right)^{-\tau_1} \left( \frac{R_0}{2(R_0)^{\frac{1}{2}}} \right)^{-s(\gamma_2 - \gamma_1)} \left( \frac{R_0^{\frac{1}{2}} + R^{\frac{1}{2}}}{R_1^{\frac{1}{2}}} \right)^{-s(\gamma_3 - \gamma_2)} \left( \frac{R_1^{\frac{1}{2}}}{R^{\frac{1}{2}}} \right)^{-s(\gamma_4 - \gamma_3)} \left( \frac{R}{R_0} \right)^{-\tau_2}, \quad (5)$$

where $\gamma_i$ are the indices of the power-laws.
where \( R_0, R_1 \) are the two break positions, \( s, s_1 \) the smoothing factors, and \( \gamma_i \) \( (i = 1, 2, 3) \) the slopes in the various rigidity ranges in between the breaks. The normalization is such that \( q_R(R) = 1 \) at \( R = R_0 \). Typically, only one break has been considered in the literature, with value of the order \( \sim 10 \text{GV} \) [9], or none \(^1\) [11]. On the other hand the recent discovery of a break at around 300 GV in the proton and helium spectra first by PAMELA [1] and then by AMS-02 [2, 3] makes it necessary to introduce a second break for a proper description of the data. This was, indeed, considered, for example, in [10, 25]. We further introduced in Eq. (5), as a novel feature with respect to previous studies, the parameters \( s_i \) to explore the possibility of a smooth transition between the various regimes, as opposed to a sharp one.

We mention here that an alternative possibility would be to model the break as a break in the diffusion rather than the injection spectrum. This has the same effect for the primaries’ spectra but leads to different results for secondaries. The secondaries’ injection spectra would reflect the break from the primary spectra, but the amount of secondaries' injection spectra would reflect the break from the primary spectra, but the amount of secondaries is typically quite low with respect to the primaries, and this allows one to evaluate Eq. (7) with a flattening above \( r \gtrsim 10 \text{kpc} \) and a cutoff above \( r \gtrsim 30 \text{kpc} \). We checked that changing the source distribution to those values has a negligible impact on the CR energy spectra after propagation.

In the case of secondary CRs, as for antiprotons produced by primary CRs through spallation in the interstellar medium (ISM), the source term is given by the primaries themselves. More precisely the source term is the integral over the momentum-dependent production rate of the secondaries and the sum over the primary species \( i \) and the ISM components \( j \),

\[
q(x, p) = \sum_{j=\text{H,He}} n_j(x) \sum_{i=\text{p,He}} \int dp_i \frac{d\sigma_{ij}(p_i, p)}{dp} \beta_i c \psi_i(x, p_i),
\]

where \( \sigma_{ij} \) is the antiproton production cross section by the species \( i \) spallating over the ISM species \( j \). The ISM is assumed to be composed of hydrogen and helium gas with fixed proportion 1:0.11. The abundance of secondaries is typically quite low with respect to the primaries, and this allows one to evaluate Eq. (7) with \( \psi_i(x, p_i) \) calculated from Eq. (1) neglecting in the first place the secondaries. Besides antiprotons, we will consider also secondary protons, i.e., primary protons that underwent inelastic scattering, losing a substantial fraction of their energy, and thus reappearing at low energies. We will also take into account tertiary antiprotons produced by the spallation of the secondary antiprotons during propagation. Secondary protons and tertiary antiprotons are described with the same formalism. Their source term can be calculated analogously to Eq. (7) but replacing \( \psi_i(x, p_i) \) with the density of primary protons in the first case, and secondary antiprotons in the second case, and using the associated production cross section. The latter is approximated as the total inelastic non annihilating cross section of the incoming proton or antiproton times the energy distribution of the scattered particle, approximated as \( 1/E_{\text{kin}} \). For more details see Ref. [29].

To numerically solve the propagation equation Eq. (1) and to derive the secondaries’ and secondaries’ abundances we use the GALPROP code\(^2\) [30, 31]. We use version r2766\(^3\) as basis, and we implement some custom modifications, such as the possibility to use species-dependent injection spectra, which is not allowed by default in GALPROP. Furthermore, we allow for a smoothing of the originally simple broken power law as discussed above.

The propagation equation Eq. (1) is solved on a grid in the energy dimension and in the two spatial dimensions \( r \) and \( z \), assuming cylindrical symmetry of our Galaxy. The radial boundary of the Galaxy is fixed to 20 kpc, while the half-height \( z_h \) is a free parameter. The radial and \( z \) grid steps are chosen as \( \Delta r = 1 \text{kpc} \) and \( \Delta z = 0.2 \text{kpc} \). The grid in kinetic energy per nucleon is logarithmic between 1 and \( 10^7 \text{MeV} \) with a step factor of 1.4. Free escape boundary conditions are used, imposing \( \psi_i \) equal to zero outside the region sampled by the grid. We tested also more accurate choices for the above settings and found the results stable against the changes.

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\(^1\) In [13, 24] a source term \( q \propto \beta^{-1}R^{-\gamma} \) is considered, which implies a break in momentum at a rigidity \( \sim m/Z \), with an upward steepening of 1 in the slope.

\(^2\) http://galprop.stanford.edu/

\(^3\) https://sourceforge.net/projects/galprop/
Note also that we consider propagation of nuclei only up to $Z=2$, i.e., in practice, in GALPROP we propagate $p$, $\bar{p}$, $^2\text{H}$, $^3\text{He}$, and $^4\text{He}$ species plus the secondary protons and the tertiaries antiprotons. This also means that we neglect possible contributions from the fragmentation of $Z>2$ nuclei, which should be a good approximation since their fluxes are much lower than the $p$ and He fluxes. Nonetheless, in the specific case of our best-fit propagation scenario (see below), we verified explicitly that including nuclei with $Z>2$ in the calculation changes the spectra of He (i.e., $^3\text{He} + ^4\text{He}$) only by few percent and protons (i.e., $p + ^2\text{H}$) by less than 1%. This is also confirmed by the study in ref.[32], where it is also shown that the $Z>2$ nuclei contribution to He is few % (although the contribution to $^2\text{H}$, $^3\text{He}$ can be, instead, up to 20-30%).

II. METHODS AND DATA

A. Data

As described in the introduction, the main focus of the analysis is on the new AMS-02 measurements. We will thus use the published proton [2] and helium [3] AMS-02 spectra, and the available preliminary measurements of the antiproton over proton ratio [14]. The AMS-02 $p$ and He data extend up to a few TV. We thus complement them with $p$ and He CREAM measurements starting from a few TV up to $\sim 100$ TV. Finally, we use recently measured $p$ and He VOYAGER data [15] at low rigidities $\lesssim 1$ GV which are believed to be the first direct measurement of the local interstellar (LIS) flux, as a consequence of the fact that the probe crossed the solar Helio pause, leaving the solar system and entering in the interstellar space. A summary of the data sets used is presented in TABLE I. We use all data in rigidity, since this is the directly measured quantity by AMS-02 as opposed to the kinetic energy.

B. Solar modulation

To compare the AMS-02 fluxes to the GALPROP model predictions solar effects have to be taken into account. CRs are deflected and decelerated in the solar winds, whose activity varies in a 22 year cycle. The effect of this solar modulation [34] can be described phenomenologically by the force-field approximation [35, 36], which is equivalent to taking into account only the adiabatic energy losses of the CRs propagating in the expanding solar wind. The process can be described by a single parameter $\phi$, the solar modulation or Fisk potential [37], which links the total energy of the particles in the local interstellar space $E_{\text{LIS}}$ to the energy $E$ observed in the detector at the Earth. The energy-differential flux $\Phi_E$ is then modulated by

$$E = E_{\text{LIS}} - |Z|e\phi$$

$$\Phi_E(E) = \frac{E^2 - m^2}{E_{\text{LIS}}^2 - m^2} \Phi_{E,\text{LIS}}(E_{\text{LIS}}),$$

where $Z$ is the charge number, $e$ the elementary charge, and $m$ the mass. The modulation potential can be approximately derived from measurements of the neutron flux at Earth by various neutron monitor stations, since a strong anticorrelation is observed between the neutron flux and the solar activity [35, 38]. Nonetheless, the procedure is affected by large uncertainties. In previous works, the usual procedure was to use this value, associate with it a “reasonable” uncertainty, and use it as a prior in the fit to CR data. Here, instead, we will use a novel procedure, similar to the one implemented in [39]. Assuming the measured VOYAGER $p$ and He fluxes, indeed, sample the LIS fluxes, we fit them with the unmodulated spectra, while, at the same time, the modulated spectra are fitted to the

| Experiment | Species | Rigidity range [GV] | ref. |
|------------|---------|---------------------|-----|
| AMS-02     | Proton  | $1.0 \cdot 10^0 - 1.8 \cdot 10^3$ | [2] |
| AMS-02     | Helium  | $1.9 \cdot 10^0 - 3.0 \cdot 10^3$ | [3] |
| AMS-02     | Antiproton ratio | $1.0 \cdot 10^0 - 0.2 \cdot 10^3$ | [14] |
| CREAM      | Proton  | $3.2 \cdot 10^1 - 2.0 \cdot 10^5$ | [33] |
| CREAM      | Helium  | $1.6 \cdot 10^1 - 1.0 \cdot 10^5$ | [33] |
| VOYAGER    | Proton  | $0.7 \cdot 10^0 - 1.0 \cdot 10^9$ | [15] |
| VOYAGER    | Helium  | $0.6 \cdot 10^0 - 2.3 \cdot 10^9$ | [15] |
AMS-02 data. CREAM data, instead, are at very high energies where solar effects can be neglected. The explicitly used $\chi^2$ is reported in the next section in Eq. (10). We thus do not assume any specific prior for $\phi$ (in practice allowing a very large range, see Table 2), and we let the VOYAGER data constrain it. In the future, monthly or weekly $p$, He, and $\bar{p}$ data from AMS-02 should further help in better constraining $\phi$.

The force-field approximation, nonetheless, gives only a first-order description of the solar modulation process. A more complete description relies on a transport equation analogous to Eq. (1) but including the specific processes experienced by CRs while propagating in the solar magnetosphere [40]. The implementation of these models is, however, beyond the scope of the present work. Indeed, dedicated analyses, using time dependent proton flux of PAMELA [41, 42] and VOYAGER data, suggest a strong rigidity and charge sign dependency of $\phi$ below a rigidity 5 GV [39, 43], indicating, in other words, a breakdown of the force-field approximation. Therefore, in the present analysis we use a fiducial lower rigidity threshold of 5 GV, although, we will also compare our fiducial fit results with those obtained including data down to 1 GV.

As a final comment, we note that while Eq. (8) is non linear, only a single average potential $\phi$ is used in it. On the other hand, $\phi$ typically undergoes significant variations during the entire period over which the final averaged measured spectrum is provided (see [44] for a recent study of the time variation of $\phi$). We thus tested a fictitious case in which $\phi$ varies linearly in time from a value of 300 MV to 700 MV during a period of 2 years and we applied to a given model LIS spectrum the force-field approximation in small time bins, using in each bin the appropriate potential and then averaging at the end to derive the final modulated spectrum. This was compared with the flux obtained by direct application of Eq. (8) to the LIS using the average $\phi$. We found no appreciable difference between the two, indicating that Eq. (8) behaves linearly to a very good approximation. A posteriori, this can be justified in terms of the smallness of the $\phi$ parameter with respect to the larger rigidities involved.

C. Fit procedure

To scan the large parameter space we use the MultiNest package [45]. MultiNest implements the algorithm of ellipsoidal nested sampling [46], allowing efficient likelihood evaluation and evidence calculation. As likelihood we use $L = \exp(-\chi^2/2)$ with

$$
\chi^2 = \frac{D}{\sigma_{AMS}} \left( \Phi_{AMS,i} - \Phi_{M}(R_i) \right)^2 + \sum_{D} \frac{\sigma_{D,i} \sum \left( \Phi_{D,i} - \Phi_{M,LIS}(R_i) \right)^2}{\sigma_{D,i}},
$$

where $D = V, C$ indicates the VOYAGER and CREAM data sets with data points $i$ at rigidity and flux $(R_i, \Phi_{D,i})$, with measured flux uncertainties flux $\sigma_{D,i}$, and $\Phi_{M,LIS}$ is the unmodulated model differential flux. Analogously, $\Phi_{AMS,i}$ and $\sigma_{AMS,i}$ indicates the AMS-02 data points and uncertainties, and $\Phi_{M}$ is the model flux modulated with a potential appropriate for AMS-02, $\phi_{AMS}$. We note here that reported fluxes by AMS-02 are actually the sum over the isotopes (protons and deuterons, and $^3$He and $^4$He). The model spectra used in the $\chi^2$ are thus also summed over the isotopes for consistency. In the above $\chi^2$ we make the simplifying assumption that all the data points are uncorrelated. This is unlikely, since in most of the rigidity range the AMS-02 errors are dominated by systematic uncertainties rather than statistical ones, and systematic uncertainties are correlated in energy in various ways. A more rigorous treatment would require a deeper knowledge of the various systematic uncertainties and a way to model them, which, however, requires a detector-level analysis not available to us. We, nonetheless, note that neglecting the correlations should imply larger errors in the CR parameters estimated from the fit, and it should thus correspond typically to a conservative assumption.

The scan is intrinsically Bayesian, in the sense that MultiNest explores the posterior, which is specified by the likelihood $L$ and the priors of the fitted parameters. Nonetheless, if the posterior and its tails have been sampled accurately enough, the likelihood samples collected by MultiNest can be also used for a frequentist analysis. This typically requires more aggressive MultiNest settings for a more accurate sampling of the likelihood. We will use as default the frequentist interpretation of the scan in terms of the profile likelihood [47], providing a comparison with the Bayesian interpretation in a specific case. As default, each scan is performed using for the MultiNest settings 400 live points, an enlargement factor $eff = 0.6$ and a tolerance $tol = 0.1$. We verified that the results are stable varying these settings. For each of the fits that we will describe in the following the typical number of likelihood evaluations performed by MultiNest is about 150,000. At the same time, with the settings described in SECTION I a GALPROP run requires about 45 CPU-seconds, for a final total computational resources usage of about $\sim$3 CPU-months. The final efficiency (number of accepted steps over computed ones) of a typical scan is found to be $\sim$7%.

In the following, contour plots for two-dimensional profile likelihoods will be shown at the 1, 2, and 3 $\sigma$ confidence levels calculated from a two-dimensional $\chi^2$ distribution. The error in the single parameters will be calculated from the related one-dimensional profile likelihood and will be quoted at 1 $\sigma$ confidence level, i.e., from the condition $\Delta\chi^2 = 1$ with respect to the minimum $\chi^2$. We will also show 1 and 2 $\sigma$ error bands around the best-fit spectra.
They are derived from the envelope spectra of all the models lying within the 1 and 2 \( \sigma \) best-fit region in the full multidimensional parameter space.

In total we perform fits with up to 11 parameters, which can be grouped in two categories. The first one includes the parameters of the shape of the injection spectrum: \( \gamma_1, \gamma_2, \gamma_1, \gamma_2, R_0, \) and \( s \). They denote the spectral indices, respectively, for protons and for the heavier species below and above the break at \( R_0 \) with smoothing \( s \) (cf. Eq. (5)). As shown in the next section, the freedom in the individual spectral indices for protons, denoted with the subscript \( p \), is necessary to achieve a good description of the measured data. We provide a dedicated study investigating the limits of a possibly universal injection spectrum at the beginning of the next SECTION III. The second category includes the parameters constraining the propagation, namely, the normalization \( D_0 \) and the slope \( \delta \) of the diffusion coefficient, the Alfvén velocity \( v_A \) related to reacceleration, the convection velocity \( v_{0,c} \), and the halo size \( z_h \). The above parameters are nonlinear and a new GALPROP run has to be performed for every new parameter set. On the other hand the fit includes three more parameters that do not require a new GALPROP run (for fixed values of the previous nonlinear parameters), namely, \( A_p, A_{He}, \) and \( \phi_{AMS} \), i.e., the normalization of the proton and helium fluxes as well as the solar modulation potential of AMS-02. For short we will call these parameters the linear parameters, even though \( \phi_{AMS} \) does not act exactly linearly. In principle, these parameters can be treated in the same way as the other, and this would give a 14-dimensional parameter space to explore. We can, however, exploit the fact that they do not require a GALPROP evaluation to simplify the problem. We, thus, do not include these parameters in the set of parameters scanned by MultiNest, but, instead, we marginalize them on the fly for each set of the other 11 MultiNest parameters. More precisely, for each nonlinear parameter set sampled by MultiNest we search for the minimum \( \chi^2 \) over the linear parameters, and we use this value to calculate the MultiNest likelihood. In this way, we use MultiNest to effectively scan over the 11-dimensional space which would be obtained from the 14-dimensional one marginalizing over the three linear parameters.

Any deviation of the normalizations \( A_i \) from 1 implies a preference for a change in the CR species abundance with respect to the input value. As the normalization and spectrum of secondaries are calculated from the primaries’ input relative abundances, rather than the ones rescaled by \( A_i \), the input abundances need to be adjusted if the best fit prefers values of \( A_i \) significantly different from 1. We thus adjust the input abundances iteratively for all fits including antiprotons, repeating the fit until the normalizations \( A_i \) converge to 1. In practice, since the initial abundances are already very close to the ones preferred by the fit, only 1 or 2 iterations are typically required for convergence. To this purpose, the output proton spectrum from GALPROP is normalized to a value of \( 4.4 \times 10^{-9} \text{cm}^{-2} \text{s}^{-1} \text{sr}^{-1} \text{MeV}^{-1} \) at a kinetic energy of 100 GeV, and the parameter \( A_p \) is thus relative to this value. The helium (\(^{4}\text{He}\)) spectrum is instead normalized to a final (i.e., found after the iterations) input abundance of \( 7.8 \times 10^4 \) relative to a proton abundance of \( 1.06 \times 10^6 \). The parameter \( A_{He} \) is thus relative to this normalization. This value was found to be appropriate (i.e., giving \( A_{He} \) compatible with 1 after the fit) for all the fits performed, except for the case uni-PHePbar (see next section) where we used an input normalization of \( 9.48 \times 10^4 \).

Three further parameters, required to specify the CR model spectra, are kept fixed or varied as a function of the

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**TABLE II: List of MultiNest and linear parameters in the fit and respective ranges of variation.** See the text for a detailed description of the parameters.

| MultiNest parameters | Ranges |
|---------------------|--------|
| \( \gamma_1 \)      | 1.2 - 2.3 |
| \( \gamma_2 \)      | 2.0 - 2.9 |
| \( \gamma_{1,p} \)  | 1.2 - 2.3 |
| \( \gamma_{2,p} \)  | 2.0 - 2.9 |
| \( R_0 \) [GV]      | 1.0 - 50 |
| \( s \)             | 0.05 - 1.0 |
| \( \delta \)        | 0.1 - 0.9 |
| \( D_0 \) [\( 10^{28} \text{cm}^2/\text{s} \)] | 0.5 - 10.0 |
| \( v_A \) [km/s]    | 0 - 60 |
| \( v_{0,c} \) [km/s] | 0 - 100 |
| \( z_h \) [kpc]     | 2 - 7 |

| Linear parameters   | Ranges |
|---------------------|--------|
| \( A_p \)           | 0.1 - 5.0 |
| \( A_{He} \)        | 0.1 - 5.0 |
| \( \phi_{AMS} \) [GV] | 0 - 1.8 |
other parameters. Specifically, the smoothness transition parameter \(s_1\) for the second break \(R_1\) is kept fixed to a value of 0.05, given the sharp transition in this case, as can be seen directly in the \(p\) and \(\text{He}\) spectra. The break itself and index after the break are fixed to \(R_1 = 450\,\text{GeV}\) and \(\gamma_3 = \gamma_2 - 0.14\), for both \(p\) and \(\text{He}\). The latter two parameters have been fixed with the following procedure: At large rigidities above \(\sim 100\,\text{GV}\) the CR spectrum is approximately given by the injection spectrum steepened by \(\delta\),

\[
\Phi_s(R) \sim q_s(R) \cdot R^{-\delta} \sim \begin{cases} (R/R_1)^{-\gamma_2-\delta} & R < R_1 \\ (R/R_1)^{-\gamma_3-\delta} & \text{else} \end{cases}.
\]

Therefore it is possible to fit a broken power law directly to the data in order to determine the break position \(R_1\) and the amount of the break \(\Delta \gamma = \gamma_3 - \gamma_2\). We performed two separate fits to \(p\) only and \(\text{He}\) only data, using AMS-02 and CREAM data together. We find that both fits give compatible results for the break position and amount of break, with the values reported above. The determination of \(R_1\) and \(\Delta \gamma\) is where CREAM data play the main role in the analysis. For the general fits performed in the following, the weight of CREAM data is quite low, since the error bars are very large compared to AMS-02. Indeed, we verified that excluding CREAM data from the fit did not significantly change the fit constraints. We also tested the case in which we exclude the CREAM data from the fit and for consistency we use values of \(R_1\) and \(\Delta \gamma\) determined from AMS-02 only. In this case the same procedure described above gives \(R_1 = 270\,\text{GV}\) and \(\Delta \gamma = -0.1\). We find that the impact of these changes on the secondary antiproton spectrum is only at the level of 10-20\% of the error bars of the data points above 50\,\text{GV}. The fit constraints, consequently, are also not significantly affected.

The fit parameters, linear and nonlinear, and their explored ranges are summarized in TABLE II.

### III. RESULTS

The results are presented as follows. We first discuss in SECTION III A the possibility of fitting the data with a universal injection spectrum. In SECTION III B we then proceed to the main analysis where separate spectral indices for the proton injection spectrum are allowed. In SECTION III C we discuss the impact of the uncertainties related to the antiproton production cross section. A comparison between the frequentist and Bayesian results is shown in SECTION III D. Finally, in SECTION III E and SECTION III F we check the robustness of the analysis results with respect to the chosen rigidity fit range and to the inclusion or exclusion of convection in the fit.

#### A. Universal injection spectrum

One of the unexpected features revealed by the precise measurements of PAMELA and AMS-02 is a significant discrepancy of the proton and helium spectral indices above \(\sim 30\,\text{GV}\), with \(\Delta \gamma_{p,\text{He}} = 0.101 \pm 0.014\) (PAMELA, [1]) and \(\Delta \gamma_{p,\text{He}} = 0.077 \pm 0.007\) (AMS-02, [2, 3]). From the theoretical point of view the reason for this difference is unclear and various possibilities have been discussed [4, 5, 7, 48–50]. Moreover, acceleration in the sources above \(\sim 30\,\text{GV}\) is expected to be charge independent and therefore the same universal injection index is expected for \(p\) and \(\text{He}\), as well as for the other species. We thus first investigate the possibility to fit the data assuming a universal injection index, attributing the difference in the observed indices to propagation effects. More in detail, we perform the fit as described in SECTION II, but we force the injection spectrum of protons and helium to be equal, i.e. \(\gamma_{1,p} = \gamma_1\) and \(\gamma_{2,p} = \gamma_2\), thus reducing the parameter space from 11 to 9 dimensions.

The fit is performed with two different data sets: in one case using only protons and helium (fit labeled as uni-PHe), and in the second case using proton, helium, and the antiproton-to-proton ratio (fit labeled uni-PHePbar). Results are shown in FIG. 1. In the (uni-PHe) case we obtain a good fit with a minimal \(\chi^2/\text{NDF}\) of 53.1 for a number of degrees of freedom (NDF) of 124. It can be seen in FIG. 1(c) that the best fit residuals with respect to the proton data are very flat in the fitted rigidity range. A similar result is obtained for the helium spectrum (not shown). The difference in the index between \(p\) and \(\text{He}\) is explained by a significant production of secondary protons that soften the observed total (primaries plus secondaries) proton spectrum by the required \(\sim 0.1\) value, with respect to the helium spectrum. In turn this imposes strong constraints in the diffusion parameter space, as can be seen by the red contours in FIG. 1(a).

In particular, a low value of \(\delta \sim 0.15\) and a low amount of reacceleration \(v_A \sim 0\) are required. Although this scenario is appealing, it is ultimately revealed to be problematic. A first problem is the amount of solar modulation required by the fit, given by \(\phi_{\text{AMS}} = 300^{+75}_{-90}\,\text{MV}\), which is quite low with respect to the neutron monitor expectation\(^4\).

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\(^4\) An updated table for the solar modulation potential up to 2016 is available under [http://cosmicrays.oulu.fi/phi/Phi_mon.txt](http://cosmicrays.oulu.fi/phi/Phi_mon.txt).
∼ 500-600 MV [38]. The second, more severe, problem is the fact that antiprotons are significantly overpredicted with respect to the observations, as shown in the lower panel of FIG. 1 (b). This can also be seen from the result of the (uni-PHePbar) fit. In fact, the parameter space constraints from this fit, shown by the black-gray contours in FIG. 1 (a), select a much larger value of $\delta \sim 0.4$ and of $v_A \sim 30$ km/s, which are incompatible, at high significance with the (uni-PHe) results. With the higher $\delta$ antiprotons data are now correctly produced but the amount of secondary protons is not enough anymore to explain the proton-helium index difference. This can clearly be seen from the systematic behavior of the residuals in FIG. 1 (d), despite the fact that, formally, the fit is still reasonable, with a $\chi^2$/NDF of 140.4/147. The (uni-PHePbar) fit also provides a more reasonable amount of solar modulation, with

(a) Triangle plot for a selected set of propagation parameters.  
(b) Predicted antiproton spectrum for the (uni-PHe) fit.  
(c) Best fit results for protons for (uni-PHe).  
(d) Best fit results for protons for (uni-PHePbar).  

FIG. 1: Comparison of the fit to data set (uni-PHe) and (uni-PHePbar) in the framework of a universal injection spectrum. In (c) and (d) the light-gray shaded regions indicate the rigidity range used in the fit. The lower panels in each of the plots show the residuals with respect to the best fit.
\[ \phi_{\text{AMS}} = 780^{+80}_{-40} \text{ MV} \]. These results are in qualitative agreement with [4, 7], where He spallation effects were studied to explain the difference in slope between proton and He, and similar difficulties in explaining secondaries spectra were encountered. Note also that, as explained in SECTION I, we are neglecting the eventual contribution of secondary protons from \( Z > 2 \) nuclei. This is not expected to be crucial, since, even in the case this contribution would be large (20-30\%) this would not solve the above issues.

In the light of the above results, we will adopt in the following as the main scenario the one in which the proton and helium spectral indices are varied independently.

**B. Main fit**

Using the 11-dimensional setup discussed in SECTION II, we will now perform fits to different data sets, to test the self-consistency of the results. In particular, we consider the following 3 fits: using only proton data (P), proton and helium data (PHe), and proton, helium and antiproton data (main). FIG. 2 shows how the propagation parameter space successively shrinks by going from data set (P) to (main). As expected, because of the large degeneracy of the parameters in case (P) nearly the whole sampled parameter space is allowed. Adding helium data results in a tight constraints on the rigidity dependence of diffusion \( \delta \sim 0.3 \), while the usual degeneracy in \( D_\nu - z_h \) appears, and no constraints on \( z_h \) can be inferred. This is also expected since strong constraints on \( z_h \) can be achieved only using precise data on radioactive clocks like \(^{10}\text{Be}, ^{9}\text{Be} \), which are not yet available. Finally, \( v_A \) and \( v_{0,c} \) are poorly constrained individually apart from a tendency to not prefer strong reacceleration \( v_A \lesssim 30 \text{ km/s} \) and a favor for large convection \( v_{0,c} \gtrsim 50 \text{ km/s} \). This is mainly due to the fact that they have approximately degenerate effects on the spectra, and only a combination of the two parameter is well constrained. Interestingly, a fit with only convection and no reacceleration seems, thus, possible. Indeed, some critical view on reacceleration has been recently discussed [51]. It can be seen that (PHe) and (main) are not compatible at the 2\( \sigma \) level, although they became fully compatible at the 3\( \sigma \) level. Given the very small error bars of AMS-02 it is perhaps expected that incompatibilities at the 2\( \sigma \) level might appear, due to the fact that the level of complexity of the fitted models is likely starting to be not comparably

**FIG. 2:** Comparison of fit results for the three data sets (P), (PHe), and (main) in the main fit framework (11 parameters) for a selected set of propagation parameters.
FIG. 3: Full triangle plot for the results of the main fit using protons, helium and antiprotons (main).

FIG. 4: Profiles of the linear parameters for the (main) fit.
FIG. 5: Comparison between data and best-fit model for the main fit framework (main): 11 parameters and fit to proton, helium, and antiprotons.

FIG. 6: Comparison between $p/p$ data and best-fit model for the main fit (main) framework (11 parameters and fit to proton, helium, and antiprotons). The various contributions to the total antiproton spectrum are also shown, as well as different production cross section models. The proton contribution includes only $pp$ production, whereas the helium contribution also includes $pHe$ and $He p$ production.

FIG. 7: $\bar{p}/p$ ratio for fit using the antiproton production cross section from di Mauro et al. [16].

adequate. We thus deem the compatibility at $3\sigma$ reasonable.

For the case (main) the complete parameter space is shown in FIG. 3 and the likelihood profiles of the linear parameters are given in FIG. 4. It can be seen that indeed $A_{He}$ is compatible with 1 at $1\sigma$. $A_p$ does actually differ from 1, but only at the 3% level which is much smaller than the uncertainties in the other parameters. We thus did not perform a further fit iteration, readjusting again the input proton normalization. The minimal $\chi^2/NDF$ of the
best fit point is 39.0/145. The agreement between data and model is thus very good (cf. also FIG.5 and FIG.6). There are no systematics features in the residuals of proton and helium spectra. The small residual structures in the antiproton-to-proton ratio are within the error band. The best-fit parameter values and their uncertainties are summarized in TABLE III. Finally, also the VOYAGER $p$ and He measurements are well fitted by the unmodulated model spectra, as shown in FIG.5. The best fit for the position of the break $R_0$ is compatible with 5 GV, our lower rigidity threshold, indicating that a low rigidity break is not necessary to fit the data. We will comment more on this point in SECTION IIIE where we show the results of the fit including data down to 1 GV.

The effect on the fit of the parameter $s$ introduced in this work can be inferred from FIG.3. It can be seen that, apart from the expected degeneracy with the break position, $s$ has only mild degeneracies with the other parameters. Indeed, performing explicitly a fit without $s$ (a sharp break), we found that only the ranges for $v_A$ and $v_{0,c}$ slightly change, the two parameters being in any case not well determined. The main effect of $s$ is, instead, to provide an overall better fit to the data and flatter residuals. It is unclear if the need for $s$ in the fit implies, indeed, that the injection spectra have a smooth break, or, alternatively if $s$ is compensating for a different effect, as, e.g., systematics in the modeling of the solar modulation.

In the following sections we will take (main) as baseline for further cross-checks and systematic studies.

C. Antiproton production cross section

The lack of precise measurements of the antiproton production cross section (cf. Eq. (7)) constitutes an important systematic uncertainty in the interpretation of the precisely measured fluxes [16, 52]. Detailed measurements for the antiproton production exist only for proton-proton inelastic scattering up to center of mass energies of $\sim$63 GeV\(^5\) [16]. For larger energies, or different target particles it is necessary to extrapolate and/or rescale the cross sections, leading to model dependent results. Furthermore, in the proton-proton inelastic scattering, only the antiproton production cross section is directly measured, while no measurement is available for the antineutron (which subsequently decay into antiproton) production cross section. In principle, from isospin symmetry the latter is expected to be equal to the former. On the other hand a measurement from NA49 [53] suggests that the antineutron cross section is actually slightly larger than the antiproton one. Further details are discussed in refs. [16, 54]. In this section we compare our

\(^5\) Some sparse measurement up to $\sim$200 GeV also exists [16].
default choice of \( p + p \to \bar{p} + X \) cross section given by Tan & Ng [55] as implemented in GALPROP with the more recent study in [16].

As mentioned above, in order to get the antiproton (\( \bar{p} \)) production cross section for arbitrary projectiles and targets, denoted by subscripts \( P \) and \( T \), respectively, we need a scaling from the \( pp \) collision cross section. A common approximation is to treat the projectile as a flux of \( A_P \) protons with momenta \( p_p/A_P \). The target is instead scaled according to the semiclassical model where the volume scales as \( A_T \) and the area as \( A_T^{2/3} \). For the production of antineutrons (\( \bar{n} \)), which subsequently decay into antiprotons and contribute directly to the flux, we assume a scaling of 1.3. Comprehensively, we get

\[
\frac{d\sigma_{P,T}(p_p,p_{\bar{p}})}{dp_{\bar{p}}} = A_P A_T^{2/3} \frac{d\sigma_{p,p}(p_p/A_T,p_{\bar{p}})}{dp_{\bar{p}}}
\]

\[
\frac{d\sigma_{P,T}(p_p,p_{\bar{n}})}{dp_{\bar{n}}} = 1.3 \frac{d\sigma_{p,p}(p_p,p_{\bar{n}})}{dp_{\bar{n}}}
\]

The dominant contribution of the antiproton flux comes from \( pp \) collisions (cf. FIG.6). The contribution from nonproton projectiles and/or targets plays a subdominant role, since both the interstellar helium gas contribution and the CR abundance of helium amount to only roughly 10%. The main uncertainty arising from the scaling is thus related to the antineutron cross section, since the antineutrons are produced directly in the \( pp \) collisions. The scaling described above gives similar results to the one implemented as default in GALPROP. FIG.6 shows the comparison between the total antiproton flux for our best fit model in the main fit framework using the default cross section and the one derived in [16] (their Eq. (13)). We can see that overall the cross section from [16] predicts a lower normalization of the antiproton flux by about \( \sim 20\% \). Also the shape is slightly different with a mild hardening of the flux starting at about \( \sim 20 \) GV. In the plot, for completeness, we also show the contribution from tertiary antiprotons, and the separate contributions from proton collisions and collisions involving helium. Finally, we also tested a different scaling available in GALPROP due to Simon et al. [56] applied to the two cross sections, and we find in both cases that this introduces a \( \sim 5\% \) variation with respect to the flux with the default scaling.

Intriguingly, as seen in FIG.6, the flux derived from the cross section from [16] seems to slightly better fit the observed antiproton spectrum. We thus repeat the fit using the new cross section (diMauro). FIG.7 shows the results using the parametrization from their Eq. (13). As expected the high-energy part fits better than in the (main) fit, but the shape of the low-energy tail does not exactly match the measurements leaving a similar amount of systematics in the residuals. As expected the lower normalization of the cross section compared to GALPROP is compensated by a slightly lower value of \( \delta \) which drops to 0.27, which can be seen from FIG.8. The other parameters are not changed. We also test the cross section from their Eq. (12), which leads to a similar result.

**D. Frequentist vs. Bayesian interpretation**

The results of this analysis are interpreted in the frequentist approach, whereas previous analyses as mostly done in a Bayesian framework. We thus compare the two approaches for the main fit case. In the Bayesian case we derive constraints from the posterior distribution, as opposed to the frequentist case where we use only the likelihood function. The Bayesian posterior is interpreted as probability distribution once the priors in model parameters are specified. In our case, prior ranges are as specified in TABLEII, and they are linear in all the parameters. Two-dimensional posteriors for two given parameters are derived marginalizing (i.e., integrating) the full posterior over the remaining parameters. In practice, marginalized posteriors are a natural output of the Monte Carlo based scanning technique, and integrals do not need to be performed explicitly. Bayesian contours are then derived integrating the marginalized posterior up to the specified confidence level. In FIG.9 we show the triangle plot for a selected set of parameters and compare the 1\( \sigma \) to 3\( \sigma \) frequentist contours with the equivalent Bayesian contours. It can be seen that the two approaches give compatible results, with the frequentist case being slightly more conservative. In fact the two approaches are expected to give compatible results in the limit in which the data are constraining enough, and the effect of the priors start to be subdominant. The above results indicate indeed that the results are data driven rather than prior driven, and thus robust.

**E. Fit down to 1 GV**

As explained in SECTION II we limit our fit range to \( R > 5 \) GV to reduce the effects of the solar modulation. Thus we avoid the rigidity range indicating rigidity and charge sign dependence of the solar modulation potential.
In this section we investigate the effect of extending the fit range down to 1 GV (labeled (1GV)). The results of the fit are shown in FIG. 10 and FIG. 11. FIG. 11 shows that a good fit is achieved with flat residuals all over the fitted energy range. The $\chi^2$/NDF has a value of 70.3/175. From FIG. 10(a) it can be seen that the two fits give consistent results at the level of a bit more than 2$\sigma$. The slight shift of about 0.04 in the value of $\delta$ from $0.28^{+0.03}_{-0.01}$ to $0.32^{+0.03}_{-0.02}$ can be considered as an estimate of the systematic error on this parameter. Regarding reacceleration and convection, adding data down to 1 GV contributes to basically break their degeneracy providing a strong constraint.
on $v_A = 25.0^{+0.92}_{-2.30}$ km/s and a lower value of $v_{0,c} = 44.4^{+12.2}_{-10.3}$ km/s. At the same time tight constraints on the index below the break $\gamma_1 = 1.32^{+0.06}_{-0.10}$ and on the break itself $R_0 = 5.52^{+0.33}_{-0.83}$ GV appear. The latter value suspiciously coincides with the rigidity below which the constant solar modulation potential approximation should start to fail. For this reason it is unclear if the presence of the break is indeed physical or if it is a way for the fit to compensate for the nonprecise solar modulation modeling. At the same time it is equally unclear if the resulting values of $v_A$ and $v_{0,c}$ are robust or are biased by the possibly incorrect solar modulation. As mentioned also in SECTION. III.B, to settle the issue a more careful study of the solar modulation effect will be necessary, complemented by the use of time series of CR data.

FIG. 11: Comparison between data and best-fit model for the global analysis including proton, helium, and antiprotons down to 1 GV (1GV).

FIG. 12: Comparison of the propagation parameters from the baseline analysis (main) to a diffusion model without convection (noVc-5GV).

FIG. 13: Comparison of the propagation parameters in the case of no convection for a rigidity cutoff at 1 GV (noVc-1GV) and 5 GV (noVc-5GV).
As last cross-check we investigate the necessity of convection. The baseline fit allows convection velocities and finds a tendency toward large values $\gtrsim 50 \text{ km/s}$, while the fit including data down to 1GV prefers low values. Therefore, we also test a diffusion model without convection (labeled (noVc-5GV)) and compare it to our baseline fit. This also allows a more direct comparison with previous works where convection was not considered [9, 10]. Results are shown in FIG. 12 and FIG. 14. The residuals in FIG. 14 show a flat behavior. The resulting $\chi^2$/NDF is 48.7/146. On the other hand the amount of solar modulation $340^{+50}_{-25}$ MV appears to be too low. FIG. 12 shows the comparison of the remaining propagation parameters to the baseline fit. The model without convection prefers slightly higher values for $D_0$, and a well constrained value of $\nu_A$, so that, overall the results seem more in agreement with the baseline fit extending down to 1GV, rather than the main case down to 5GV. Given also the low value for the solar modulation in the case (noVc-5GV) we thus also tested the case of data down to 1GV (labeled noVc-1GV). In this case the $\chi^2$/NDF increases to 82.2/176, but we achieve a more reasonable value for the solar modulation potential of $640 \pm 20$ MV. The propagation parameters are not much affected by the data below 5GV as can be seen from FIG. 13. FIG. 15 shows that residuals are reasonably flat also for this fit.
We report in TABLE III a summary of the 1σ constraints on the parameters for the various fits performed. When the lower or upper range coincides with the chosen prior, the constraint is reported in italized characters. The χ² values for each fit, also broken into the sub-data set, used are reported in TABLE IV. We can use the results from TABLE III to derive the systematic uncertainties on δ. Averaging between the (main) fit and the (1GV) fit we get a value of δ = 0.30±0.03. From the fits without convection we see that δ can be upshifted by up to a value of 0.1, while in the (diMauro) fits δ can be downshifted by a value of 0.04. We thus quote these last two numbers as systematic uncertainties so that δ = 0.30±0.03(stat)+0.10(sys).

### IV. SUMMARY AND CONCLUSION

We have presented new constraints on the propagation of Galactic CRs from an (up to) 11-dimensional parameter fit to the latest AMS-02 spectra for p, He, and p/p. Solar modulation is treated within the force-field approximation, but the modulation potential is constrained with a novel approach, fitting the unmodulated CR p and He spectra to recently available low-energy data from VOYAGER, collected after the probe left the heliosphere and thus sampling...
the local interstellar CR flux. The VOYAGER data and the unmodulated spectra are fitted jointly to the AMS-02 data and the modulated spectra.

As a first attempt, we try to fit the data with a universal injection spectrum for $p$ and He. We find that a universal injection is possible when fitting only $p$ and He data. In this case, the observed difference in $p$ and He slopes of about $\sim 0.1$ can be explained by a significant production of secondary $p$ so that the total primary plus secondary $p$ spectrum is steepened by the required 0.1 value in the slope. However, this requires a quite low value of the spectral index of diffusion $\delta \sim 0.15$, and implies a large production of $\bar{p}$ which significantly overpredicts the observations. This scenario is, thus, in the final instance, not viable. For the main results we thus perform a fit leaving individual spectral freedom to $p$ and He. With this additional freedom a good fit to $p$, He, $\bar{p}/p$ spectra is achieved. The main result is a tight constraint on $\delta = 0.30_{-0.02}^{+0.03} (\text{stat})_{-0.10}^{+0.10} (\text{sys})$, where the error is dominated by systematic uncertainties rather than statistical ones. The robustness of this result has been cross-checked against various factors, like the uncertainties in the solar modulation, the choice of the diffusion model framework, i.e., if convection is allowed or not, and the systematic uncertainties in the $\bar{p}$ production cross section. Since solar modulation is most important at low energies, its effect was studied using different cuts (1 and 5 GV) on the AMS-02 data. The effect of uncertainties in the $\bar{p}$ production cross section was, instead, tested comparing the results of the fit when different available determinations of the cross section are used. Both of these effects have an order $10 - 20\%$ impact on the value of $\delta$, while the most important effect is the inclusion of convection in the model, which shifts the value of $\delta$ from $\sim 0.3$ to $\sim 0.4$.

For the other propagation parameters the results are less definitive. The height of the Galactic halo and the normalization of diffusion present a well-known degeneracy, which, not surprisingly, cannot be resolved. In this respect, more precise “CR-clocks” measurements, like the ratio $^{9}\text{Be}/^{10}\text{Be}$, which will be available in the future from AMS-02, are necessary. Regarding convection and reacceleration, the fit above 5 GV prefers large convection velocities of $v_{0,c} \gtrsim 50\text{ km/s}$ and Alfven velocities of $v_{A} \lesssim 25\text{ km/s}$, with large parameter errors coming from a degeneracy between convection and reacceleration. The fit with data down to 1 GV breaks this degeneracy and gives a well definite reacceleration of $v_{A} = 25 \pm 2\text{ km/s}$ and preference for lower values of $v_{0,c} \lesssim 50\text{ km/s}$. It remains, however, unclear how robust this determination of $v_{A}$ is, since it relies on data below 5 GV which are significantly affected by solar modulation. Finally, we find that a fit without convection is nonetheless possible, providing a good fit to the $p$, He and $\bar{p}/p$ data, and giving a similar value of $v_{A}$.

A comparison of these results from the constraints imposed from the AMS-02 observations of lithium, boron, and carbon will be presented in a forthcoming companion paper.

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