On the structure of the turbulent interstellar atomic hydrogen

I. Physical characteristics.
Influence and nature of turbulence in a thermally bistable flow

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Aims. We study in some details the statistical properties of the turbulent 2-phase interstellar atomic gas.

Methods. We present high resolution bidimensional numerical simulations of the interstellar atomic hydrogen which describe it over 3 to 4 orders of magnitude in spatial scales.

Results. The simulations produce naturally small scale structures having either large or small column density. It is tempting to propose that the former are connected to the tiny small scale structures observed in the ISM. We compute the mass spectrum of CNM structures and find that \( N(M)dM \propto M^{-1.7}dM \), which is remarkably similar to the mass spectrum inferred for the CO clumps. We propose a theoretical explanation based on a formalism inspired from the Press & Schecter (1974, ApJ, 187, 425) approach and use the fact that the turbulence within WNM is subsonic. This theory predicts \( N(M) \propto M^{-5/3} \) in 2D and \( N(M) \propto M^{-16/9} \) in 3D. We compute the velocity and the density power-spectra and conclude that, although the latter is rather flat, as observed in supersonic isothermal simulations, the former follows the Kolmogorov prediction and is dominated by its solenoidal component. This is due to the bistable nature of the flow which produces large density fluctuations, even when the rms Mach number (of WNM) is not large. We also find that, whereas the energy at large scales is mainly in the WNM, at smaller scales, it is dominated by the kinetic energy of the CNM fragments.

Conclusions. We find that turbulence in a thermally bistable flow like the atomic interstellar hydrogen, is somehow different from turbulence in a supersonic isothermal gas. In a companion paper, we compare the numerical results with atomic hydrogen observations and show that the simulations well reproduce various observational features.

Key words. hydrodynamics – instabilities – ISM: kinematics and dynamics – ISM: structure – ISM: clouds

1. Introduction

The neutral atomic gas (HI) is ubiquitous in the galaxies and is of great importance for the star formation process, since molecular clouds form by contraction of HI. As such, HI has been extensively observed over the years (e.g. Kulkarni & Heiles 1987; Dickey & Lockman 1990; Heiles & Troland 2003, 2005; Miville-Deschênes et al. 2003). Although HI has also received a lot of attention from the theoretical point of view (Field 1965; Field et al. 1969; Zel’dovich & Pikel’ner 1969; Penston & Brown 1970; Wolfire et al. 1995, 2003) and in spite of the early recognition that HI is a turbulent medium (e.g. Crovisier 1981; Heiles & Troland 2005), it is only recently that the dynamical properties of HI have been investigated.

Two lines have been simultaneously pursued by various teams. Attempting to understand the properties of HI on large scales, Gazol et al. (2001, 2005), Dib & Burkert (2005), de Avillez & Breitschwerdt (2005), have performed 2D or 3D numerical simulations of turbulent multi-phase flows subject to various forcing. On smaller scales, various studies attempting to resolve the physical scales involved in the problem, have focused on the description of the thermal contraction, trying to understand the cooling and the condensation of a piece of WNM into CNM accurately. Hennebelle & Pérault (1999, 2000) have studied the influence of a converging flow, Koyama & Inutsuka (2000) have investigated the influence of a shock propagating in a bistable flow whereas Sánchez-Salcedo et al. (2002) have considered initially unstable gas. More recently, Hennebelle & Passot (2006) have investigated the influence of Alfvén waves propagating into the medium. Because of the high numerical resolution required to treat the problem, these works were performed in only one dimension. Further important extensions of these works have been performed in 2 or 3D. Koyama & Inutsuka (2002) have generalized their study in 2D and Kritsuk & Norman (2002) have computed the evolution of thermally unstable gas using 3D simulations.

In a recent paper (Audit & Hennebelle 2005, here after Paper I), we have investigated the dynamical evolution of colliding flows of WNM, paying particular attention to the effect of turbulence. The main conclusions of this study are as follows: first, we confirm, in a context where the flow is strongly turbulent, the result obtained by Koyama & Inutsuka (2002) in their study of shock propagation in WNM. The CNM is very fragmented in small clouds which have a dispersion velocity equal to a fraction of the WNM sound speed. Second, as in Gazol et al. (2001), although at much smaller scales, we find large fractions...
of thermally unstable gas and we show that turbulence is able to stabilize transiently the thermally unstable gas. Third, we show that in spite of the strong turbulence, the CNM fragments are generally not very far from pressure equilibrium with the surrounding WNM. Colliding flows of atomic interstellar gas have been further investigated by Heitsch et al. (2005, 2006) and by Vázquez-Semadeni et al. (2006).

One of the crucial issues in modeling atomic hydrogen is the numerical resolution which is very demanding (see Sect. 2.3). In this paper, we present the result of very high resolution numerical simulations allowing description of scales ten times smaller than in Paper I. We perform a set of simulations with various numerical resolutions to study its influence. We also study the influence of thermal conduction by varying its value. The high resolution allows a better description of the flow down to much smaller scales and offers the possibility to perform various statistical studies of the flow and CNM structures properties.

In a companion paper (Hennebelle et al. 2007, Paper III), we make preliminary comparisons between the 2D simulations presented here and various observational results. In particular, the high resolution permits the comparison with various small scale structures that have been observed in HI, like the so-called tiny small atomic structure (TSAS) and recent low column density clouds observed by Braun & Kanekar (2005) and Stanimirović & Heiles (2005).

In Sect. 2, we describe the numerical setup and method and discuss briefly the various scales which have to be adequately treated. In Sect. 3, we present one typical timestep of the simulation and discuss qualitatively various features of the flow which are worth quantifying. Section 4 presents the density and the pressure PDF. Section 5 presents density and velocity power-spectra as well as the energy spectrum. In Sect. 6, the mass spectrum of the CNM structures in the simulation is computed and in Sect. 7, a theoretical explanation is proposed. Section 8 concludes the paper.

2. Initial conditions and method

The initial conditions and the methods are very similar to those of Paper I. Here, we describe them briefly for self-consistency.

2.1. Equations, notations and numerical method

We consider the usual fluid equations for a radiatively cooling gas including thermal conductivity namely,

\[ \partial_t \rho + \nabla \cdot [\rho u] = 0, \]

\[ \partial_t \rho u + \nabla \cdot [\rho u u + P] = 0, \]

\[ \partial_t \rho u + \nabla \cdot [\rho u (E + P)] = -\nabla \cdot (L(\rho, T)) + \nabla (\kappa(T) \nabla T). \] (3)

\[ \rho \] is the mass density, \( u \) the velocity, \( P \) the pressure, \( E \) the total energy and \( L \) the cooling function which includes Lyman-\( \alpha \), \( C^+ \) and \( O \) line cooling and grains photodissociation heating. The gas is assumed to be a perfect gas with \( \gamma = 5/3 \) and with a mean molecular weight \( \mu = 1.4 m_H \), where \( m_H \) is the mass of the proton. \( \kappa \) is the thermal conductivity and is given by \( \kappa(T) = \gamma C_T \rho(T) \) where \( C_T = k_B / m_H(\gamma - 1) \), \( \gamma = 5.7 \times 10^{-3} \) (T/1 K)\(^{1/2} \) g cm\(^{-1} \) s\(^{-1} \) and \( k_B \) is the Boltzmann constant.

We use the HERACLES code to perform the simulation. This is a second order Godunov type hydrodynamical code. Godunov methods are now widely used and tested and are particularly well suited to the treatment of shocks (e.g. Toro 1997). The size of the computational domain is 20 pc and the resolution ranges from 600\(^2 \) to 10 000\(^2 \) cells leading to a spatial resolution ranging from 3.3 \times 10^{-2} \ pc to 2 \times 10^{-2} \ pc.

2.2. Initial and boundary conditions

The boundary conditions consist in an imposed converging flow at the left and right faces of amplitude 1.5 \times C_s \_wnm on top of which turbulent fluctuations of amplitude \( \epsilon = 2 \) (see Paper I for accurate definitions) have been superimposed. On top and bottom faces, outflow conditions have been setup. This means that the flow is free to escape the computational box across these 2 faces.

The initial conditions consist of uniform WNM (\( n \approx 0.8 \) cm\(^{-3} \)) at thermal equilibrium. It is worth stressing that initially no CNM is present in the computational box. The simulations are then runned until a statistically stationary state is reached. Typically, this requires about 5 to 10 box crossing times. At this stage, the incoming flow compensates on average the outgoing material.

For the highest resolution runs, this would lead to very long simulations and we therefore adopt a slightly different strategy. We start the run with a lower resolution of 1250\(^2 \) cells and wait until the statistically stationary regime is obtained. Then, we increase the resolution of the simulation by a factor 2 and restart the now 4 times larger simulation. This new simulation is runned until the stationary regime is reached. Typically this takes about two box crossing time. This procedure is then repeated until the desired spatial resolution is obtained.

2.3. Scales

Contrary to polytropic fluids, a 2-phase medium has various spatial and temporal characteristic scales which have to be adequately described. Here, we briefly recall them.

2.3.1. A static spatial scale

The Field length (Field 1965) is the length at which thermal diffusivity becomes comparable to the heating and cooling terms. Its typical value is about 0.1 pc in the WNM and about 10\(^2 \) pc in the CNM. The Field length is also the typical scale of the thermal fronts that connects the cold and warm phases. As demonstrated by Field (1965), it determines the smallest wavelength at which thermal instability can grow and, therefore, the size of the smallest CNM structures. The effect of varying the thermal diffusivity is discussed in the following.

2.3.2. Two dynamical scales

We distinguish three scales whose origin is due to dynamical processes. The first scale is the cooling length of the WNM, i.e. the product of the cooling time and the sound speed within WNM, \( \lambda_{\text{cool}} = \tau_{\text{cool}} \times C_s \_\text{wnm} \). This scale is about 10 pc and corresponds to the typical length at which WNM is non linearly unstable and can be dynamically triggered by a compression into the unstable regime.

Since the ratio between the CNM and WNM density is about hundred, the size of a CNM structure formed through the contraction of a piece of WNM of size \( \lambda_{\text{cool}} \), will be typically, hundred time smaller (assuming a monodimensional compression). Therefore, the size of the CNM structures is about
cooling length of WNM divided by hundred, leading to about $\lambda_{\text{cool}}/100 \approx 0.1 \text{ pc}$.

Finally, as first pointed out by Koyama & Inutsuka (2002), the fragments of CNM have a velocity dispersion equal to a fraction of the sound speed of the medium in which they are embedded, i.e. WNM. Since the contrast between the sound speed within the two phases is about 10, it means that CNM structures undergo collisions at Mach number, $M \approx 10$. If, for simplicity, we assume isothermality and apply Rankine-Hugoniot relations, we obtain that the size of the shocked CNM structures is given by the size of the CNM structures divided by $M^2$ which is about $10^{-2} \text{ pc}$, whereas the density peak is given by $100 \times M^2 \approx 10^3 \text{ cm}^{-3}$. Note that since the effective polytropic index of the CNM, $\gamma_{\text{eff}}$, is closer to 0.7 than to 1, these numbers are underestimated by a factor of about 7.

The consequences of not solving properly these scales have been previously discussed qualitatively in Paper I. In the following, quantitative estimates are given.

### 2.4. Comparison with other simulations

As explained in the previous section, it is actually not possible to treat simultaneously all the scales involved in the physics of the atomic gas. This implies compromises and induces various choices of initial, boundary and forcing conditions. In order to make clear the domain of validity of various related works, we now discuss the numerical and physical setups which have been chosen. It is worth stressing that the different choices are complementary and equally interesting since they allow one to focus on different aspects of the same physical problem.

Koyama & Inutsuka (2002) have investigated the propagation of a shock into WNM. They have an extremely high spatial resolution ($7 \times 10^{-4} \text{ pc}$) and a relatively modest box size of about 0.25 pc (in the transverse direction). With these choices, the CNM structures are very well resolved but the statistics are low and the structures are small. In a recent paper (Koyama & Inutsuka 2006), they consider a larger box and try to understand the properties of the turbulence that develops without any external forcing.

Heitsch et al. (2005, 2006) and Vázquez-Semadeni et al. (2006) have focused on the formation of a CNM turbulent layer. They consider a simulation box of about $10 \text{ pc}$ and have a resolution of about $10^{-5} \text{ pc}$. Such a setup which is very similar to the one used in Paper I, allows to treat reasonably well the scales, but does not provide a description of the CNM structures as accurate as the CNM structures obtained in Koyama & Inutsuka (2002). Also in these work, the authors focus on the transient regime, rather than on a time-independent situation.

Gazol et al. (2001, 2005) and Piontek & Ostriker (2004) have focused on larger simulation boxes (100 pc–1 kpc) and coarser spatial resolution (0.5–5 pc). The same kind of scales has been investigated by de Avillez & Breitschwerdt (2005) using an AMR scheme. This leads to a much better description of the larger scales and a more realistic self-consistent forcing (e.g. due to supernovae or the development of the magneto-rotational instability) that the choice makes in this work, but only marginal description of CNM structures can be achieved.

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1 In fact, as we will see in the following, the size of the CNM structures follows a density distribution. The numbers given here are simply indicative.
are produced by supersonic collisions between CNM fragments. It should be noted that the density peak is not fully resolved because of insufficient numerical resolution and therefore it is expected that this event should have indeed a larger peak density and a smaller size if the numerical resolution was higher. It is also important to emphasize that the cooling function used in this simulation is probably not accurate for such high densities.

3.3. Small column density clouds

In a similar way, the properties of the smallest structures that form in the simulations appear to be reminiscent of the tiny structures which have been recently observed by Braun & Kanekar (2005) and Stanimirović & Heiles (2005). Many of them can be seen in Fig. 2 (e.g. for $y \leq 6$ pc) and a cut through two small CNM structures is displayed in Fig. 5 ($y \approx 6.68$ and 6.83 pc). The density of these two structures is about 30 and 100 cm$^{-3}$ whereas their size is about $10^{-2}$ pc.

Further examples are shown in Paper III in which synthetic HI spectra are also discussed.

3.4. Pressure distribution

Another important aspect is revealed by the pressure field displayed in Fig. 3. The pressure field appears well organized with large scale systematic gradients. At large scales, the pressure and the density fields appear to be broadly anti-correlated. Indeed, the central cloud (located roughly between $x = 1$ and $x = 4$ and $y = 6$ and $y = 9$) is surrounded by high pressure WNM, whereas its internal pressure is a few times lower. This is a consequence of the radiative cooling. When the WNM fluid particles enter into the shocked layer, their internal pressure goes up like in an adiabatic shock, then they cool and finally condense out. Indeed, the cooling length of shocked (Mach 1.5–2) WNM is few parsec which is the length of the high pressure WNM layer seen on the left part of the box. Interestingly, the high density shocked CNM is located at the interface between the high pressure WNM and the low pressure cloud. Note that since the WNM pressure ($P \approx 10^4$ K cm$^{-3}$) is lower by at least one order of magnitude than the large pressure fluctuation ($P \approx 10^5$ K cm$^{-3}$), it cannot be the triggering agent. However, this high pressure WNM is in part responsible of these large pressure fluctuations since it actively induces the formation of relatively high velocity CNM structures, which then undergo supersonic collisions.

At small scales, the situation is much different. The pressure is poorly correlated with the density field displayed in Fig. 2. Most of the CNM structures do not have high thermal pressure (see Fig. 5) with the notable exception of the densest regions which correspond to shocked CNM. Indeed, some of the CNM structures have a thermal pressure lower than the surrounding WNM as for example, the 2 structures located at $y = 7.15$ and 7.3 pc in Fig. 5. This low pressure indicates that the structures have just formed, so that there was not enough time for the initially low thermal pressure induced by the radiative cooling to readjust to the surrounding higher pressure.

Finally, the density field indicates that the CNM structures are well identified and connected to the surrounding WNM by very stiff fronts. Moreover, as in the static 2-phase medium (Field et al. 1969; Wolfire et al. 1995), most of the CNM structures are not far from quasi-pressure equilibrium with the surrounding medium (in the sense that pressure fluctuations are small with respect to the density contrast between the two phases). This is likely a consequence of the large...
density contrast between the phases. First, with a density contrast of about hundred, the sonic waves propagating in the WNM as well as the eddies get mostly reflected when they reach a WNM/CNM interface, probably making the energy injection inside the structures relatively inefficient. Second, since the scale of the CNM structures is small compared to the scale of the WNM flow and since most of the turbulent energy is on the largest scales, the WNM flow is relatively uniform at the scale of the CNM structures and mostly advects them uniformly.

To summarize, it seems that a turbulent 2-phase flow cannot be accurately described as a polytropic turbulent flow, neither as a static 2-phase medium, but, instead, they present a kind of
duality, some aspects being more reminiscent of turbulent flows and other of static 2-phase media with fluctuations which can be up to hundred times the mean CNM density or pressure. In the following, we therefore quantify the mass distribution of CNM structures, as well as the density and pressure distribution of the flow. We also study the statistics of the flow by computing various power-spectra, in order to characterize its scale dependence. In Paper III, we quantify further the numerical results by computing line of sight statistics and various properties of the CNM structures.

4. Density and pressure probability distribution function

Figures 6 and 7 show the probability distribution function of the logarithm of the density and pressure respectively. Full black lines correspond to the runs having \(l_g^2 = 10000^2\) cells, whereas the dotted red lines are for the runs with \(l_g^2 = 5000^2\) cells. The density pdf presents two peaks which correspond to the WNM and CNM densities. The pdf values at densities between 5 and 50 indicates that there is a significant fraction of thermally unstable gas, as first stated by Gazol et al. (2001) and confirmed in Paper I.

In principle, the density pdf allows estimation of the fraction of the dense gas that has been compressed by the ram pressure.
However it is seen that the results for the high density part do clearly depend on the numerical resolution, meaning that higher resolution runs will lead to a higher fraction of high density gas. It should also be clear that since the fraction of high density gas depends on the cooling processes, it will certainly be modified by a more accurate treatment of the microphysics. Having these limitations in mind, we can indicatively say that the fraction of gas denser than $10^3 \, \text{cm}^{-3}$ is about 1–3% for this simulation. It is important to keep in mind that, as emphasized in Paper I and in Gazol et al. (2005), this obviously depends on the driving of the turbulence.

The pressure pdf indicates that the average pressure is about $6000 \, \text{K cm}^{-3}$, which is the pressure of the WNM injected at the boundary. The large pressures are induced by the ram pressure of the incoming flow. The pressure distribution is reminiscent of pdf found by various authors in different contexts (Passot & Vázquez-Semadeni 1998; Scalo et al. 1998). Comparison with the results presented by Gazol et al. (2005) is more straightforward since they consider a thermally bistable flow as well. As in their case, we find that a near-powerlaw tail develops at high and low pressure (in their case they report that this depends on the driving length). Because of the higher numerical resolution, we reach higher values of pressure. Our pressure pdf appears to be somehow similar to the results presented in their Fig. 11 corresponding to a large scale driving. However, the significant difference between the 2 numerical experiments precludes detailed comparisons.

5. Velocity and density power-spectra and energy spectrum

5.1. Incompressible and isothermal turbulence

An important diagnostic commonly used in fluid dynamics to characterize the scale-dependent structure of the flow is the power-spectrum of the fluid variables such as the velocity and the density fields. The power-spectrum equal to the square of the power-spectrum of the fluid variables such as the velocity and the density fields. The power-spectrum of pdf found by various authors in different contexts (Passot & Vázquez-Semadeni 1998; Scalo et al. 1998). Comparison with the results presented by Gazol et al. (2005) is more straightforward since they consider a thermally bistable flow as well. As in their case, we find that a near-powerlaw tail develops at high and low pressure (in their case they report that this depends on the driving length). Because of the higher numerical resolution, we reach higher values of pressure. Our pressure pdf appears to be somehow similar to the results presented in their Fig. 11 corresponding to a large scale driving. However, the significant difference between the 2 numerical experiments precludes detailed comparisons.

In incompressible turbulence, the velocity power-spectrum, $P_V$, is related to the energy spectrum as $E(k) \propto k^{D-1} P_V$, where $D$ is the space dimension. The Kolmogorov (Kolmogorov 1941) prediction for the energy spectrum of homogeneous, 3D and incompressible turbulence is: $E(k) \propto k^{-5/3}$. Note that this corresponds to power-spectrum proportional to $k^{11/3}$ for 3D flows and to $P_V \propto k^{-8/3}$ for 2D flows. However, the behaviour of 2D incompressible turbulence is very different from the 3D turbulence because of enstrophy conservation (Kraichnan 1967). At scales larger than the injection scale, one expects to recover the Kolmogorov energy spectrum whereas at scales smaller than the injection scale, Kraichnan (1967) predicts $k P_V \propto k^{-8}$.

In 1D compressible turbulence, when shocks dominate, a Burgers-like spectrum $P_V \propto k^{-2}$ is expected (Elsasser & Schamel 1976). Passot et al. (1988) have confirmed, using isothermal bidimensional numerical simulations, that the compressible part of the velocity field follows the same behaviour whereas for the solenoidal part, the prediction of Kraichnan (1967) is well verified, $k P_V \propto k^{-3}$. Kim & Ryu (2005) recently explored numerically in some details the 3D case. They show that the density power-spectrum ($P_\rho$) for low rms Mach number flow is close to the Kolmogorov power-spectrum, whereas it is much flatter at high Mach Mach number, i.e. $k^2 P_\rho \propto k^{-\alpha}$ with $\alpha = 0.5$ for an rms Mach equal to 12. They interpret this result as the consequence of the density distribution being highly concentrated in sheets and filaments. Flat density power-spectra have also been observed in MHD isothermal simulations. Padoan et al. (2004) report $\alpha = 0.25$ for a model with equipartition between magnetic and kinetic energy and $\alpha = 0.75$ for a super-Alfvénic model (see also Beresnyak et al. 2005).

5.2. Velocity power-spectrum

Figure 8 shows the velocity power-spectrum for various numerical resolutions, whereas Fig. 9 shows the compensated power-spectra (i.e. $k^2 P_V$ the value of $a$ being given in the figure) of the solenoidal and compressible components of the velocity field. For reference the Kolmogorov (1941) and the Kraichnan (1967) prediction have been displayed in Fig. 8. As in Passot et al. (1988), the compressible part has a power-spectrum $k P_V \propto k^{-2}$. The velocity power-spectrum, as well as the power-spectrum of the solenoidal component, appear to be in good agreement with the Kolmogorov law, $k P_V \propto k^{-5/3}$ for $k$ between $\approx -3.3$ and $\approx 1.8$.

This may be at first surprising because, as recalled above, the prediction of Kraichnan for 2D incompressible fluid is that there is an inert cascade above the energy injection scale, leading to the velocity power-spectrum predicted by Kolmogorov and a direct enstrophy cascade at scale below the injection scale leading to $k P_V \propto k^{-3}$. However, this result is a consequence of the conservation of vorticity in incompressible or barotropic flow. When the flow is non barotropic the vorticity equation writes:

$$\partial_t \omega + \nabla \times (\omega \times v) = \nabla \times \nabla \left( \frac{1}{\rho} \right),$$

(4)

In a radiatively cooling flow, as the interstellar atomic hydrogen, the pressure and the density are not necessarily correlated as already emphasized in Sect. 3. This means that the baroclinic
term on the right hand side, is not small with respect to the second term of the left hand side, implying that enstrophy is not conserved. To verify this, we have computed the mean value of the (norm of the) baroclinic term and of the second lhs term over the whole simulation. We find that the former is on average 10 times smaller than the latter. We also find that for about 7% of our grid points, the ratio between the two is higher than 0.5. It seems likely to us that this constitutes a sufficiently large deviation from the strict enstrophy conservation to explain the deviation from Kraichnan’s prediction. Unsurprisingly, we find that the higher values of the baroclinic term are obtained in the fronts where the density changes much but not the pressure.

To confirm the influence of the baroclinic term, we have performed isothermal simulations with exactly the same setup as simulations with cooling (same code and same initial and boundary conditions). In these simulations, the baroclinic term vanishes exactly. The compensated spectra are shown in Fig. 10. The power-spectrum of the solenoidal component is very stiff and roughly proportional to \( k^{-4.4} \) whereas the compressible component has a power-spectrum proportional to about \( k^{-3.2} \). These numbers are indeed much closer to the results of Passot et al. (1988) and to the prediction of Kraichnan (1967).

Finally, we note that, as revealed by Fig. 9, although compressible and shear modes are comparable at large scales, the second dominates at small scales for \( \log(k) > -3 \) (solid and dotted lines are very close between \(-1 \geq \log(k) > -3\)).

5.3. Density power-spectrum

Figure 11 shows the power-spectrum of the density field. It appears to be flat since one finds \( k_\rho \propto k^{-0.4} \). This is reminiscent of what is observed by Kim & Ryu (2005) although in the present case the strong and stiff density fluctuations are induced by the 2-phase nature of the flow rather than by highly supersonic motions.

Kim & Ryu (2005) proposed that the flat density power-spectrum can be understood by the fact that the Fourier transform of a Dirac function is independent of \( k \) leading to \( P_\rho \propto k^0 \) in 1D. We believe that this explanation is also valid in the present case and that the density power-spectrum is related to the size distribution of the structures (Figs. 15–17). Indeed since the structures are bounded by contact discontinuities, they have stiff boundaries and they resemble, depending on the scales at which one is considering them, either to the Dirac or Heaviside functions.

As a matter of fact, the end of the inertial range which occurs at about \( k = 0.1 \) (about 10 cells in the simulation) corresponds to the length at which the structure distribution is influenced by the numerical diffusivity implying that the number of density peaks at these scales becomes gradually smaller.

Interestingly, Desphande et al. (2000) using 21 cm line absorption data probing linear scales between \( 10^{-2} \) and 3 pc, therefore comparable to the present simulation, report density power-spectrum \( k^2 P_\rho \propto k^{-\alpha} \), in two different regions for which \( \alpha = 0.75 \) and \( \alpha = 0.5 \). These values are significantly flatter than the Kolmogorov index and at least for the second one, are very close to the value of 0.4 which is inferred in our simulations.

5.4. Energy spectrum

Figure 12 shows the power-spectrum, \( P_E \), of the quantity \( \sqrt{E} \). The quantity \( k^2 P_\rho \) corresponds to the energy spectrum in the incompressible case and following Klessen et al. (2000), we call it the energy spectrum, \( E(k) \). It is rather different from the Kolmogorov prediction and significantly flatter. In the inertial range \( E(k) \propto k^{-0.9} \), meaning that the energy is approximately equally distributed in the \( k \)-space. This appears to be a direct consequence of the density power-spectrum discussed above which relies on the strong density fluctuations due to the 2-phase
nature of the flow. In order to confirm this, we have calculated the energy spectrum and the power-spectrum of the square root of \( \rho \) for data obtained by taking \( \min(\rho, \rho_0) \). The results for various values of \( \rho_0 \), namely 2, 3 and 10 cm\(^{-3} \) as well as the result for the untruncated field, are shown in Figs. 13 and 14. When \( \rho_0 \) decreases, \( P_0^\rho \) becomes stiffer (note that for the untruncated field, we find \( k P_0^\rho \propto k^{-0.6} \) in the inertial range). The same behaviour is also observed for the energy spectrum which becomes closer to the velocity power-spectrum shown in Fig. 8.

For \( k \) larger than \( \approx 5 \times 10^{-3} \) (corresponding to a scale of about 0.5 pc) and \( \rho_0 < 3 \text{ cm}^{-3} \), the energy spectrum becomes significantly lower than the energy spectrum obtained for the untruncated field. This clearly indicates that at scales smaller than about 0.5 pc, most of the energy is stored in the CNM bulk motions. In other terms, in our numerical experiment, the 2-phase nature of the flow appears to play an important rôle and constitutes a significant deviation of single phase hydrodynamics at scales smaller or comparable to about 1 pc.

6. Mass spectrum

As in Paper I, we extract the structures by a simple clipping algorithm. Since the cold structures are connected to the surrounding warm gas by stiff thermal fronts such simple criteria has a clear physical meaning. The threshold is equal to 30 cm\(^{-3} \) which lies in the thermally unstable regime (note that we have verified that taking 10 cm\(^{-3} \) instead does not change the results significantly).

6.1. Numerical result

Figure 15 shows the number of structures, \( n_s \), per logarithmic mass interval, \( \delta n_s/d \log M \) (note that since the simulations are 2D, the mass is expressed in solar mass per parsec). The largest structures have a mass of about one solar mass per parsec whereas the mass of the smallest structures is about \( \approx 10^{-5} \) solar mass per parsec. The number of structures, \( N(M) \), decreases with mass, and follows approximately: \( \delta n_s/d \log M \propto 1/M^{\beta-1} \) with \( \beta \approx 1.7 \), until \( \log(M) \approx -3 \). This implies that the number of structures per mass interval is \( \delta n_s/dM = N(M) \propto 1/M^{\beta} \).

For \( \log(M) \leq -3 \), \( \delta n_s/d \log M \) increases with mass. This part of the distribution is most likely affected by the numerical resolution. This is confirmed by the fact that the numbers of low mass objects decreases with the numerical resolution. Interestingly enough, the number of massive structures becomes independent of the resolution for \( dx \leq 10^{-3} \) pc (see solid, dotted and dashed curves). This indicates that numerical convergence is reached for the large structures (log\( (M) \geq 3 \)). It is however clear that no numerical convergence has yet been reached for the smallest structures. We also note that the number of structures more massive than \( \approx 0.3 \text{ M}_\odot \text{ pc}^{-1} \), appears to be smaller than the value expected from the power-law fitted above. This is most likely due to the finite size of the numerical experiment.

For \( \log(M) \geq -3 \), the total mass contained in a logarithmic interval of mass, follows approximately \( dM_{tot}/d \log M \propto M^\beta N \approx M^{2-\beta} \approx 0.25 \text{ M}_\odot \text{ pc}^{-1} \), which indicates that, although the mass of CNM is, in principle, dominated by the large clouds, the index of the power-law is nevertheless very shallow. This is confirmed by Fig. 16 which shows \( dM_{tot}/d \log M \) as a function of \( \log(M) \). It is seen that the mass in large structures is only slightly larger than the mass in hundred to thousand times smaller structures. This indicates that the description of the small structures is very important in order to achieve a fair description of the flow and its dynamics.

Complementary information is given in Fig. 17 which shows the total number of cells, \( \delta n_{tot}/d \log n_c \) per logarithmic interval of computational cells number, \( n_c \), contained in the structure (expressed in cells of the highest resolution run). The physical size of the structures is obtained by taking the square root of the cell number and multiplying by \( 2 \times 10^{-3} \) pc which is the numerical resolution of the 10000\(^\circ \) cells run. The size at which the structure number starts to decrease because of insufficient numerical resolution is about 10 (\( \approx \sqrt{n_c} \) for \( n_c \approx 100 \)). The length
All runs have been performed with a resolution of 5000^2 cells. We therefore have to introduce a lower value in scales. We thus performed various runs in which its value has been changed. In order to investigate the exact influence of the thermal conductivity on the structures, we have pointed out by Koyama & Inutsuka (2004). They show explicit expressions for the probability of finding a contrast density \( \delta = \rho/\langle \rho \rangle - 1 \), \( \langle \rho \rangle \) being the mean density, is

\[
P_\delta(\delta) = \frac{1}{\sqrt{2\pi}\sigma(R)} \exp\left(-\frac{\delta^2}{2\sigma(R)^2}\right)
\]

(5)

where

\[
\sigma^2(R) = \int \delta^2(k)W_k^2(R)d^3k
\]

(6)

(see e.g. Padmanabhan 1993, Chap. 5). In this last expression \( \delta(k) \) is the power-spectrum of \( \delta \).

Therefore, the first question that must be addressed is whether this assumption is justified in the case of the atomic hydrogen. More precisely, since the CNM structures, are the final product of density fluctuations arising in the WNM, the question which has to be addressed is whether the spectrum of the density fluctuations in the WNM, smoothed at a scale \( R \), is reasonably described by the probability stated by Eq. (5) and what is the value of \( \sigma(R) \). One difficulty to address this question is that in the multiphase simulations presented previously, the WNM and the CNM are mixed together. They are therefore not representative of the initial state of the WNM which leads to the CNM structures. On the other hand, the early times of the simulation reflect the numerical setup rather than the initial conditions. To overcome this difficulty, we consider the simulation with cooling presented above but in which we have removed the CNM by taking \( \min(\rho, p_0) \), with \( p_0 = 3 \) cm^{-3} (see Sect. 5.4). As shown by Fig. 13, the power-spectrum of \( \sqrt{\theta} \) is nearly Kolmogorov. We have checked that the power-spectrum of the density field is indeed nearly Kolmogorov as well. Another point is that the power-spectrum of the density in the isothermal simulation performed in Sect. 5.2 (let us remember that this simulation has exactly the same initial, boundary and forcing conditions than the other runs performed with cooling) is indeed very close to the Kolmogorov spectrum. The reason is that the isothermal simulation is subsonic (as for the simulation with cooling, the rms velocity is about half the sound speed of the WNM). Since in the ISM, the WNM is likely to be more or less isothermal until the thermal pressure is triggered to a value for which WNM is thermally unstable, this should constitute a reasonable description of WNM before the thermal transition occurs. Therefore, we assume that \( \tilde{\delta}^2(k) \propto k^{-n} \), with \( n \approx 8/3 \).

To calculate the width of the distribution of the density fluctuations for the case of the turbulent WNM, we choose the simple window function sharply truncated in \( k \)-space, \( W_\ell(R) = \theta(R^\ell - k) \) where \( \theta(z) = 1 \) if \( z \geq 0 \) and 0 otherwise. With Eq. (6), it is easy to see that the integral diverges for small \( k \). This is a natural consequence of the well known property of the Kolmogorov spectrum that most of the energy is on the large scales. We therefore have to introduce a lower value in \( k \)-space,
which corresponds to an upper limit of the spatial scales, $L_c$. In the case of the simulation, this is naturally given by the size of the computational box whereas in the ISM, this would be the energy injection scale. We have:

$$\sigma^2(R) = \int_{|\delta_c|}^{R}|L_c| \delta^2 \frac{d\delta}{\delta_c} = C \frac{1}{n-2} \left( \left( \frac{1}{L_c} \right)^{2-n} - \left( \frac{1}{R} \right)^{2-n} \right), \quad (7)$$

where $C$ is a constant.

The question is then, are the density fluctuations reasonably described by Eqs. (5) and (7)? To answer this question, we have smoothed the density field of the isothermal and cooling simulations using the simple window function, $W_b(\rho_c) = 1/b^2 \sum_{|\delta| \leq |\delta_c|} |\rho_c - \rho|$, for various values of $b$. Figures 19 and 20 show the results in the isothermal case and in the case with cooling (since the isothermal simulation has 5000^2 cells, the values of $b$ are 2 times lower than in the case with cooling). It is seen that, in the isothermal case, the probability distribution is reasonably Gaussian. As expected from Eq. (7), $\sigma$ appears to be a decreasing function of $b = R/\delta_c$. In the case with cooling, the situation is slightly less clear because the shape is a little skewed towards the large densities. This is a consequence of the fact that a fraction of the gas is thermally unstable and in transition towards CNM or is already CNM. This dense gas is therefore on the process of condensation or already condensed and should not be considered in the pdf of density fluctuations of WNM.

Beyond this qualitative agreement, we present in Appendix A a more quantitative estimate that confirms the validity of Eq. (7). Thus it appears that the assumption of the distribution of the density fluctuations being given by Eqs. (5) and (7) is sufficiently justified.

7.2. Mass function

To calculate the mass function, we can now follow the same procedure as for the cosmological calculation. We note, however, two important differences. First, since we assume statistical stationarity, there is no time dependence in our analysis. Second, the collapse is not due to gravity but to thermal instability.

Here, since our primary goal is to understand the simulation results, we consider the 2D case. Thus, for a perturbation of size, $L$, the final mass of the structures is about $M(L) = \pi L^2$, Eq. (7) is equivalent to

$$\sigma^2(M) = \frac{C}{n-2} \left( M(L_c)^{(n-2)/2} - M^{(n-2)/2} \right) \frac{1}{M^{(n-2)/2}}, \quad (8)$$

where $M = M(R)$.

The fraction of objects having a mass larger than $M$ is thus given by:

$$F(M) = \int_0^\infty P_2(\delta) s(\delta) d\delta, \quad (9)$$

where $s$ is the selection function$^2$. The selection function, $s$, determines the fluctuations that will eventually lead to the formation of a CNM structure. For simplicity, we will assume that once the density reaches a threshold $\delta_c$, it leads to a thermal collapse and to the formation of a CNM structure. Although this assumption is reasonable given the physical characteristics of thermal instability, we note that various improvements could certainly be possible, as for example considering a density-pressure relation for the instability threshold or taking into account the stabilizing rôle of the turbulence within the perturbation (in that case $s$ would also depend on the velocity dispersion). Thus we have

$$F(M) = \int_0^\infty P_2(\delta) s(\delta) d\delta. \quad (10)$$

The mass function, $N(M)$, is then simply given by $-\partial F/M(M/\overline{M})$. We have$^3$

$$N(M)dM = -\frac{1}{\sqrt{2\pi} \overline{M} \sigma^2} \frac{d\sigma}{dM} \exp\left( -\frac{\delta^2}{2\sigma^2} \right) dM. \quad (11)$$

With Eq. (8), we get

$$\frac{d\sigma}{dM} = -\sqrt{\frac{Cp}{2}} \sigma \left( \frac{n}{n-2} \right) \frac{M^{(n-2)/2}}{4M} \times \left( \frac{(n-2)}{M(L_c)^{(n-2)/2} - M^{(n-2)/2}} \right). \quad (12)$$

This leads to:

$$N(M)dM \propto M^{(n-2)/2-2-1} \frac{1}{\sigma^2} \exp(-\sigma^2/(2\sigma^2))dM. \quad (13)$$

Therefore for small masses, i.e. mass smaller than $M(L_c)$, we obtain that $N(M) \propto M^{n-2}/n$ with $n \approx 8/3$, we get $N(M) \propto M^{-5/3}$. We note that $5/3 \approx 1.666$ is rather close to the value 1.7

$^2$ We note that in cosmology this expression is known to be inconsistent because in the cosmological case, the integral of $dF = f(m)dm$ from 0 to $\infty$ should be equal to 1. This problem has been solved by the Excursion set theory (Bond et al. 1991). The results for the shape of the mass spectrum is however independent of this and we will not consider this problem further at this stage.

$^3$ The Excursion set theory would give the same result multiplied by a factor 2.
obtained in the simulation. For a mass close to $M(L_c)$, there is an exponential cutoff which is due to the cut, $L_c$ introduced at large scales. Altogether these results are similar to the results obtained in the cosmological case. An important difference however lies in the exponent of the mass spectrum which is less stiff. This is a consequence of the fact that in a turbulent flow, most of the energy is at large scales implying more large scale fluctuations.

7.3. The 3D case: similarity with the CO clumps

This analysis can be straightforwardly applied to the 3D case. The only differences are that $n = 11/3$ (since the simulations presented here are bidimensional, it is worth to remember that Kim & Ryu (2005) do find that in transonic isothermal simulations, the density power spectrum is nearly Kolmogorov) and $M = \mathbb{P}R^3$. We obtain:

$$\frac{d\sigma}{dM} \propto \frac{M^{(n-3)/3}}{M} \left( \frac{(n-3)}{M(L_c)^{(n-3)/3} - M^{(n-3)/3}} \right),$$

and

$$N(M)dM \propto M^{(n-3)/3-2} \frac{1}{\alpha^3} \exp(-\delta^2 / (2\alpha^2))dM.$$

The index of the power law of the mass spectrum in the 3D case and for masses small compared to $M(L_c)$, is about $2 - (n-3)/3 = 16/9 \approx 1.78$, i.e. slightly stiffer than the value inferred in the 2D case.

Interestingly, we note that the value inferred here, turns out to be similar to the index of the mass spectrum observed for the CO clumps of masses between $10^5$ solar mass and one Jupiter (Kramer et al. 1998; Heithausen et al. 1998).

Although it cannot be excluded that this is a pure coincidence, one possibility is that the origin of the CO clumps is indeed rooted in the atomic gas. The mass distribution of the CO clumps may therefore be determined during the atomic phase before the gas becomes molecular.

8. Conclusion

We have presented high resolution numerical simulations aiming to describe a turbulent interstellar atomic flow. The high resolution ($2 \times 10^{-3}$ pc) provides a good description (although not sufficient to reach numerical convergence) of the flow close to the spatial scales of the smallest structures observed in HI. We confirm the main results obtained in Paper I. The flow is very fragmented and the phases are tightly interwoven. The CNM structures are connected to the WNM by sti ff thermal fronts and are locally in near pressure equilibrium with the surrounding WNM. Altogether, this is reminiscent of the classical 2-phase model in spite of the fact that the flow is dynamical and turbulent.

We find that small scale structures, either diffuse or very dense, are naturally produced in a turbulent 2-phase medium. Whereas the former are simply the tail of the structure mass spectrum induced by turbulence, the latter, which are somehow reminiscent of the TSAS observed in the atomic gas, are transient events produced by supersonic collisions between CNM fragments.

We characterize the flow by studying various statistical quantities, namely the mass distribution of the CNM structures, the velocity and density power-spectrum and the energy spectrum, paying special attention to the numerical resolution. We stress that the present problem requires a lot of numerical resolutions to obtain reliable results. Typically at least $5000^3$ to $10000^3$ cells are needed and even so, no strict convergence is achieved.

Our main results are as follows.

For structures of size larger than about 10 grid cells, for which numerical diffusivity is not too important, we find a mass spectrum, $N(M) \propto M^{-1.7}$. In order to explain this result, we have carried out a calculation based on Press & Schecter (1974) formalism. Our theory predicts $N(M) \propto M^{-3/2}$ in 2D and $N(M) \propto M^{-10/9}$ in 3D. One of the main assumptions is that the CNM clumps are due to density fluctuations within WNM, which behaves as a sub to transonic nearly isothermal gas. We note that these mass spectra are similar to the mass spectrum inferred observationally for CO clumps. This is compatible with the origin of CO clumps being rooted in the very diffuse atomic gas although this does not, by any means, constitute a proof.

Like in highly supersonic isothermal flows, the density power-spectrum is rather flat. However, the structures being pressure-bound ed and not bounded by shocks as in supersonic isothermal simulations, they are long-lived objects. Unlike supersonic isothermal simulations, the velocity power-spectrum follows the Kolmogorov prediction and is dominated by its solenoidal part. At scales larger than about 1 pc, most of the energy is in the WNM whereas at scales smaller than about 1 pc, it is mostly in the CNM. Indeed, the energy is nearly equally distributed in $k$-space for scales ranging between about 2 and $2 \times 10^{-2}$ pc where numerical diffusion becomes dominant. This behaviour is due to the density structure of the flow that presents relatively flat density power-spectrum, indicating that in a thermally bistable flow like the one we studied, the kinetic energy at scales below 1 pc, appears to be dominated by the translational motions of the CNM fragments.

We also study the influence of the thermal conduction by performing various runs with different thermal conductivities and find a modest influence, mainly on the small structures.

In a companion paper (Paper III), we further characterize the simulation results by studying line of sight statistics and the physical properties of the CNM clouds. We also calculate synthetic HI spectra and reach the conclusion that, although the lines of sight from which they are calculated are very complex, showing in some cases several structures, the spectra are relatively smooth and simply broadened by the complex motions along the line of sight.

Altogether these results suggest that the turbulence which takes place in the neutral atomic interstellar gas is very different from the turbulence which takes place in an isothermal or polytropic gas. One important question that remains to be investigated is how these conclusions will change in a more realistic framework of 3D flows. To answer this question 3D simulations have been performed and will be presented elsewhere (Audit & Hennebelle 2007).

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Appendix A: Further measurement of the pdf of density fluctuations

In order to measure more accurately the dependence of $\sigma$ (Eq. (7)) on $R$, we define $\delta_{2h} - \delta_{2b} = W_{2h}(p) - W_{2b}(p)$, i.e. the density field smoothed by the window function $W_{2h} - W_{2b}$. With this window function, one finds that
Fig. A.1. Isothermal simulation. Probability distribution of the density field smoothed by the window function $W_k - W_{2k}$ for various value of $b$. The range of the $x$-axis being proportional to $b^{n-2}$, it appears that the length of these distribution is reasonably Gaussian and roughly proportional to $b^{n-2}$.

Fig. A.2. Same as Fig. A.1 for the 2-phase simulation in which the density field has been truncated by taking $\min(\rho, \rho_0)$. 

$\sigma(R) = C/\sqrt{(n-2) \times (2^{n-2} -1)} R^{n-2}$. In particular, it is independent of $L$ and is proportional to $R^{n-2} = R^{2/3}$ for $n = 8/3$.

Figures A.1 and A.2 display the distribution of $\delta \rho_b - \delta \rho_2$ for various values of $b$ for the isothermal and the 2-phase simulations. The range of the $x$-axis is proportional to $b^{n-2}$. As can be seen, the distributions are reasonably Gaussian and the width is broadly proportional to $b^{n-2}$ for both cases except for small $b$ in the isothermal simulation.

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