Numerical solution of the Boltzmann equation for the shock wave in a gas mixture

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Abstract

We study the structure of a shock wave for a two-, three- and four-component gas mixture on the basis of numerical solution of the Boltzmann equation for the model of hard sphere molecules. For the evaluation of collision integrals we use the Conservative Projection Method developed by F.G. Tcheremissine which we extended to gas mixtures in cylindrical coordinates. The transition from the upstream to downstream uniform state is presented by macroscopic values and distribution functions. The obtained results were compared with numerical and experimental results of other authors.

List of symbols

$\rho_1, \rho_2$ molecular mass of $i$-th component
$p_i$ molecular momentum of $i$-th component
$x$ vector of configuration space
$f_i(p_i, x, t)$ distribution function of $i$-th component
$\bar{p}_i = \frac{p_i}{m_2} - \frac{\bar{p}_1}{m_1}$ initial relative velocity
$(p_{x,i}, p_{r,i})$ coordinates of momentum in cylindrical coordinates
$M$ Mach number
$R$ gas constant

k Boltzmann constant
$p$ nodes of momentum grids
$\Omega$ domain in momentum space
$V$ volume of domain $\Omega$
$N_0$ number of momentum nodes
$N_v$ number of integration nodes
d diameter of molecules of $i$-th component
$n_i$ number density of $i$-th component
$u_i$ flow velocity of $i$-th component
$T_i$ temperature of $i$-th component
$T_{xx,i}$ parallel temperature of $i$-th component
$T_{rr,i}$ transversal temperature of $i$-th component
$n$ number density of the mixture
$\rho$ mass density of the mixture
$u$ flow velocity of the mixture
$T$ temperature of the mixture
$l$ mean free path of molecules
$\tau$ mean collision time
$\chi_i$ concentration of $i$-th component

1 Introduction

The shock wave structure for a binary gas mixture is an important problem in kinetic theory. It has been investigated experimentally and theoretically by using moment methods, direct simulation Monte-Carlo method (DSMC), fluid dynamics methods, numerical analysis based on kinetic models, conservative splitting method, finite-difference analysis of the Boltzmann equation (see a review in [1]).

The first numerical solution of the Boltzmann equation for a single gas was obtained in [2] and then
in [3]. Later, the problem was solved by discrete-coordinate methods for Boltzmann equation with different techniques of evaluation of the collision integral: polynomial approximation of distribution function in velocity space [4–5], application of polynomial correction for fulfilling conservation laws [6], conservative projection method [7–10].

The method of [5] was extended to binary gas mixtures [11], the method of polynomial correction [6] was applied to binary gas mixtures in [12–13], the method of [8–9] was extended at first to binary gas mixtures [11] and then to three- and four-component mixtures [15]. Later computations for 3-component mixtures were repeated by the same method in [19].

In this paper the shock wave structure for two-three- and four-component gas mixtures is solved by an extension of Conservative Projection Method [7–10] applied to the complete kinetic Boltzmann equation. The method is based on a special projection techniques for evaluation of the collision integrals. The computed collision integral is conservative for density, momentum and energy. It is equal to zero when the solution has a form of Maxwellian distribution function. The integration grid for evaluation of the collision integral is given in [17]. The differential part of the Boltzmann equation is approximated by conservative finite-difference scheme of the second order [18]. In this scheme the transport of mass, momentum and energy between the nodes of the configuration space is realized in a conservative way.

Here we have studied the behavior of densities, flow velocities, parallel, transversal and total temperatures for the mixture and its components for various Mach numbers, various masses and concentration ratios. The results were compared with the results of [11] and experiments [19] with a good agreement between them.

2 Description of the method

The system of Boltzmann kinetic equations for a mixture of monatomic gases containing $K$ components is usually written in the form

$$\frac{\partial f_i}{\partial t} + \vec{v}_i \frac{\partial f_i}{\partial x} = I_i, \quad i = 1, \ldots, K. \quad (1)$$

The collision integrals have the form

$$I_i = \sum_j \int_{\mathbb{R}^3} 2\pi b_m \int_0^\infty (F'_i F'_j - F_i F_j) g d b d \varepsilon d \vec{\xi}_j. \quad (2)$$

Here we have used the notation: $F_i = F_i(\vec{\xi}_i, \vec{x}, t)$, $F'_i = F_i(\vec{\xi}'_i, \vec{x}, t)$, $\vec{\xi}_i$, $\vec{\xi}'_i$ and $\vec{\xi}', \vec{\xi}_i$ are velocity vectors before and after collisions, respectively, $g = |\vec{\xi}' - \vec{\xi}_i|$, $b_m$ is the maximum interaction distance, $b$ and $\varepsilon$ are impact parameters of a binary collision. To extend the conservative method of evaluating collision integrals [10] to gas mixtures, it is sufficient to transform equation (1) from the velocity variables to momentum variables:

$$(\vec{\xi}_i, \vec{x}, t) \mapsto (\vec{p}_i, \vec{x}, t), \quad F_i = F_i(\vec{\xi}_i, \vec{x}, t) \mapsto f_i(\vec{p}_i, \vec{x}, t).$$

From the normalization condition $\int F_i d \vec{\xi}_i = \int f_i d \vec{p}_i = n_i$, one obtains $F_i(\vec{\xi}_i, \vec{x}, t) = f_i(\vec{p}_i, \vec{x}, t)m_i^2$. The system of Boltzmann equations in the momentum space take the form

$$\frac{\partial f_i}{\partial t} + \frac{\vec{p}_i}{m_i} \frac{\partial f_i}{\partial \vec{x}} = I_i. \quad (3)$$

Collision integrals (2) become

$$I_i = \sum_j \int_{\mathbb{R}^3} 2\pi b_m \int_0^\infty (f'_i f'_j - f_i f_j) g d b d \varepsilon d \vec{p}_j, \quad (4)$$

where

$$f_i = f_i(\vec{p}_i, \vec{x}, t), \quad f'_i = f_i(\vec{p}'_i, \vec{x}, t), \quad g = \left| \frac{\vec{p}'_i}{m_2} - \frac{\vec{p}_i}{m_1} \right|. \quad (5)$$

The following properties should be conserved in a discrete form of the collision integral:

$$\int_{\mathbb{R}^3} I_i(\vec{p}_i) \Psi(\vec{p}_i) d \vec{p}_i = 0, \quad \Psi(\vec{p}_i) = \left(1, \frac{\vec{p}_i}{m_1} \right),$$

$$I_i[f_{i,M}] = 0, \quad f_{i,M} = n_i \left( \frac{1}{2\pi k T m_i} \right)^{3/2} \exp \left( -\frac{(\vec{p}_i - \vec{p}_0)^2}{m_i 2kT} \right).$$

2 Description of the method
System (3) with collision integrals (4) is solved either on a uniform 3-dimensional grid $S_0$ with $N_0$ points $p_\gamma$ in Cartesian momentum space $\Omega$ of volume $V$ or on a uniform 2-dimensional grid in cylindrical coordinate system due to the cylindrical symmetry of the problem. For brevity, values of collision integrals and distribution functions at the grid nodes are denoted by $I_{i,\gamma}$ and $f_{n,\gamma}$ ($\alpha = i,j$), respectively. The use of a constant step in the coordinate space is needed for conservation of the total momentum in the projection method.

System (3) of $K$ equations is transformed to the system of $N_0K$ equations:

$$\frac{\partial f_{i,\gamma}}{\partial t} + \frac{\partial f_{i,\gamma}}{\partial x} = I_{i,\gamma},$$

$$i = 1, 2, \ldots, K, \ \gamma = 1, 2, \ldots, N_0.$$  

Evaluation of collision integrals is performed with the use of 8-dimensional uniform integration grid $S_\nu = (\bar{p}_{i,\nu}, \bar{p}_{j,\nu}, b_{\nu}, \varepsilon_{\nu})$ in the domain $\Omega \times [0, 2\pi] \times [0, b_{\nu}]$ with $N_\nu$ nodes (the cross $\times$ denotes direct product) in such a way that momenta $\bar{p}_{i,\nu}$ and $\bar{p}_{j,\nu}$ coincide with momentum grid nodes while all the variables $b_{\nu}, \varepsilon_{\nu}$, for which the post-collision momentum $\bar{p}_{i,\nu}$ or $\bar{p}_{j,\nu}$ falls outside of $\Omega$, are excluded. The collision integral for $n$-th component at node $\gamma$ can be written in the form

$$I_{n,\gamma} = \frac{1}{4} \sum_{i,j} \int_0^{2\pi} \int_0^{b_{\nu}} \Phi_{n,\gamma}(f_{i,f_{i}} - f_{i,j}) d\varphi dx d\bar{p}_{i} d\bar{p}_{j}$$

(6)

with $\Phi_{n,\gamma}$ being the following combination of Dirac $\delta$-functions and Kroneker symbols $\delta_{n,l}$ ($\delta_{n,l} = 1$ if $n = l$ and $\delta_{n,l} = 0$ if $n \neq l$):

$$\Phi_{n,\gamma} = \delta_{n,l} \delta(\tilde{p}_i - \bar{p}_{i,\nu}) + \delta_{n,l} \delta(\tilde{p}_j - \bar{p}_{j,\nu}) - \delta_{n,l} \delta(\tilde{p}_i - \bar{p}_{i,\nu}) - \delta_{n,l} \delta(\tilde{p}_j - \bar{p}_{j,\nu}).$$

(7)

The conservative projection method for evaluation of (6) imply replacing the two last $\delta$-functions in (7) by their decompositions with a splitting coefficient $r_\nu \ll 1$ which has to be defined from the energy conservation law. For each contribution to the integral sum this decomposition has the form (omitting subscript $\nu$):

$$\delta(\tilde{p}_i' - \bar{p}_{i,\nu}) = (1 - r)\delta(\tilde{p}_i - \bar{p}_{i,\nu}) + r\delta(\tilde{p}_i, \lambda + s - \bar{p}_{i,\nu})$$

$$\delta(\tilde{p}_j' - \bar{p}_{j,\nu}) = (1 - r)\delta(\tilde{p}_j, \mu - \bar{p}_{j,\nu}) + r\delta(\tilde{p}_j, \mu - s - \bar{p}_{j,\nu}).$$

(8)

In (5) the grid nodes $\bar{p}_{i,\lambda}$ and $\bar{p}_{j,\mu}$ are the closest ones to the post-collision vectors $\tilde{p}_i'$ and $\tilde{p}_j'$, respectively, whereas $\bar{p}_{i,\lambda + s}$ and $\bar{p}_{j,\mu - s}$ are some complementary nearly located grid nodes. Hence the contributions to the collision integral in two near-grid points are replaced by the weighted contributions in two pairs of the closest nodes. A necessary condition to make this decomposition conservative is the fulfillment of the momentum conservation law. Using a uniform grid in $\Omega$ for the kinetic equation in the momentum space ensures the fulfillment of this condition.

Coefficient $r$ (subscript $\nu$ is skipped) is defined from the energy conservation law

$$E_0 = (1 - r)E_1 + rE_2, \ \text{where} \ \ E_0 = \frac{\bar{p}_{i,\lambda}^2}{2m_i} + \frac{\bar{p}_{j,\mu}^2}{2m_j},$$

$$E_1 = \frac{\bar{p}_{i,\lambda}^2}{2m_i} + \frac{\bar{p}_{j,\mu}^2}{2m_j}, \ \ E_2 = \frac{\bar{p}_{i,\lambda + s}^2}{2m_i} + \frac{\bar{p}_{j,\mu - s}^2}{2m_j}.$$  

(9)

2.1 Details of calculations in cylindrical coordinates

For computing the shock wave structure it is convenient to pass to cylindrical coordinate system $(p_x, p_r, \varphi)$ in momentum space in which the distribution function $f_i(\bar{p}_x, \bar{p}_r, \bar{p}_\varphi)$ is replaced by the function $f_i(\bar{p}_x, \bar{p}_r, \bar{p}_\varphi, x, t)$ independent of $\varphi$ due to the assumed cylindrical symmetry of the problem. In domain $\Omega$ of volume $V$ in momentum space we introduce a uniform two-dimensional grid with $N_0$ points $(p_{x,\nu}, p_{r,\nu}, \varphi)$.

Equation (5) becomes

$$\frac{\partial f_{i,\gamma}}{\partial t} + \frac{p_{x,\nu} + p_{r,\nu}}{m_i} \frac{\partial f_{i,\gamma}}{\partial x} = I_{i,\gamma},$$

$$i = 1, 2, \ldots, K, \ \gamma = 1, 2, \ldots, N_0.$$  

For the evaluation of collision integrals we use the grid

$$S_{\nu} = \{(p_{x,\nu}, p_{r,\nu}, \varphi) \in \Omega \times [0, 2\pi], \ (p_{x,\nu}, p_{r,\nu}, \varphi) \notin \Omega \times [0, 2\pi]\}$$
with \( N_e \) nodes in such a way that \((p_{x,i',\nu}, p_{r,i,\nu})\) and \((p_{x,j',\nu}, p_{r,j,\nu})\) coincide with the momentum grid nodes, while the angles are distributed uniformly in corresponding intervals. The integral (6) must be written in cylindrical coordinates:

\[
I_{n,\gamma}^c = \frac{1}{4} \sum_i \sum_j \int_0^{2\pi} \int_0^{2\pi} \int_0^b \int_0^b \Phi_{n,\gamma}(f_j', f_j^* - f_i, f_j) g \times p_{r,i} p_{r,j} d p_{r,i} d p_{r,j} b d b d \varphi d \theta d \varphi.
\]

In formulas (7) and (8) we make the following replacements:

\[
\delta(p_i - \tilde{p}_i, \gamma) = \delta(p_{x,i} - p_{x,i',\gamma}) \delta(p_{r,i} - p_{r,i,\gamma}),
\]

\[
\delta(p_j - \tilde{p}_j, \gamma) = \delta(p_{x,j} - p_{x,j',\gamma}) \delta(p_{r,j} - p_{r,j,\gamma})
\]

and

\[
\delta(p_i' - \tilde{p}_i', \gamma) = (1 - q) \delta(p_{x,i'} - p_{x,i',\gamma}) \delta(p_{r,i'} - p_{r,i',\gamma}) + q \delta(p_{x,i,\lambda + s} - p_{x,i'}, \gamma) \delta(p_{r,i,\lambda + s} - p_{r,i',\gamma}),
\]

\[
\delta(p_j' - \tilde{p}_j', \gamma) = (1 - q) \delta(p_{x,j'} - p_{x,j',\gamma}) \delta(p_{r,j'} - p_{r,j',\gamma}) + q \delta(p_{x,j,\mu - s} - p_{x,j'} - p_{r,j,\mu - s} - p_{r,j',\gamma}),
\]

respectively. Here the nodes nearest to \( \tilde{p}_i' \) and \( \tilde{p}_j' \) are \((p_{x,i,\lambda}, p_{r,i,\lambda})\) and \((p_{x,j,\mu}, p_{r,j,\mu})\), respectively. The two sets of four nodes surrounding the points \( \tilde{p}_i' \) and \( \tilde{p}_j' \) can be denoted by \((p_{x,i,\lambda + s}, p_{r,i,\lambda + s})\) and \((p_{x,j,\mu - s}, p_{r,j,\mu - s})\), where \( \vec{s} \) and \( \vec{\tilde{s}} \) are vectors of displacement along the grid. In Cartesian coordinates from the well-known relation between vectors of momentum before and after the collision we have \( \vec{s} = -\vec{\tilde{s}} \). However, in cylindrical coordinates we have only one equality for the first \( x \)-coordinate.

The decomposition coefficient \( q_{\nu} \) (\( \nu \) has been skipped in the formulas above and below) can be determined from the energy conservation law:

\[
E_0 = E_1(1 - q) + q E_2,
\]

where

\[
E_0 = \frac{p_{x,i}^2 + p_{r,i}^2}{2m_i} + \frac{p_{x,j}^2 + p_{r,j}^2}{2m_j},
\]

\[
E_1 = \frac{p_{x,i,\lambda}^2 + p_{r,i,\lambda}^2}{2m_i} + \frac{p_{x,j,\mu}^2 + p_{r,j,\mu}^2}{2m_j},
\]

\[
E_2 = \frac{p_{x,i,\lambda + s}^2 + p_{r,i,\lambda + s}^2}{2m_i} + \frac{p_{x,j,\mu - s}^2 + p_{r,j,\mu - s}^2}{2m_j}.
\]

We have demonstrated in some detail how a transformation of variables from velocity space to momentum space (in Cartesian and cylindrical coordinates) in the system of Boltzmann equations makes it possible to build the conservative projection method for evaluation of collision integrals.

After calculating the collision integral, the system of discrete ordinate equations (5) is solved by the standard procedure of applying the splitting method until the stabilization of the solution.

The obtained distribution functions define the following gas dynamics parameters for components: the number densities \( n_i \), flow velocities \( u_i \), temperatures \( T_i \), parallel temperatures \( T_{xx,i} \), and transversal (radial) temperatures \( T_{rr,i} \). For the whole gas, one obtains the molecular number density \( n \), density \( \rho \), flow velocity \( u \), and temperature \( T \). The listed macroscopic variables are defined as sums of moments of the distribution functions. For a gas mixture in cylindrical coordinates one has

\[
n_i = \sum_{\gamma} p_{r,i,\gamma} f_{i,\gamma}, \quad u_i = \frac{1}{n_i m_i} \sum_{\gamma} p_{r,i,\gamma} p_{x,i,\gamma} f_{i,\gamma},
\]

\[
T_{xx,i} = \frac{1}{k n_i m_i} \sum_{\gamma} p_{r,i,\gamma} (p_{x,i,\gamma} - u_{x,i} m_i) f_{i,\gamma},
\]

\[
T_{rr,i} = \frac{1}{2k n_i m_i} \sum_{\gamma} p_{r,i,\gamma} f_{i,\gamma},
\]

\[
T_i = \frac{1}{3} (2T_{rr,i} + T_{xx,i}), \quad n = \sum_i n_i,
\]

\[
\rho = \sum_i m_i n_i, \quad u = \frac{1}{\rho} \sum_i m_i n_i u_i,
\]

\[
T = \frac{1}{k n} \sum_i [k u_i T_i + m_i n_i (u_i - u)^2/3].
\]

3 The problem of a shock wave structure

We consider a plain shock wave traveling in \( x \) direction with Mach number \( M \). We denote the parameters: numerical densities, flow velocity and temperature before the shock wave as \( n_1^{(1)}, n_1^{(2)}, u_1^{(1)}, T_1^{(1)} \), respectively, and those behind the shock wave as \( n_2^{(2)} \),
Parameters on both sides of the shock wave are related by the Rankine-Hugoniot conditions for monatomic gas

\[
n^{(2)}_i / n^{(1)}_i = u^{(1)} / u^{(2)} = 4M^2 / (M^2 + 3),
\]

\[
T^{(2)} / T^{(1)} = (5M^2 - 1)(M^2 + 3)/16M^2,
\]

where \(k\) is the Boltzmann constant, \(m_1\) and \(m_2\) are the masses of the first and second component, respectively, \(\chi_1^{(1)}\) and \(\chi_2^{(1)}\) are concentrations of the first and second components before the shock wave. Here \(M\) is Mach number before the shock wave.

The problem is solved in the coordinate system attached to shock wave. The steady shock wave structure is obtained as the evolution of initial discontinuity of gas parameters posed at \(x = 0\). Boundary conditions are imposed at sufficiently large distances from the discontinuity at \(x = -L_1, x = L_2\), where the gas can be considered as being in thermodynamic equilibrium with the corresponding Maxwellian distribution functions:

\[
f_i(t = 0, x < 0, p_{x,i}, p_{r,i}) = f_i(t, x = -L_1, p_{x,i}, p_{r,i}) = \frac{n_i^{(1)}}{(2\pi T^{(1)} m_i)^{3/2}} \exp \left( -\frac{(p_{x,i} - u^{(1)} m_i)^2 + p_{r,i}^2}{2kT^{(1)} m_i} \right),
\]

\[
f_i(t = 0, x > 0, p_{x,i}, p_{r,i}) = f_i(t, x = L_2, p_{x,i}, p_{r,i}) = \frac{n_i^{(2)}}{(2\pi T^{(2)} m_i)^{3/2}} \exp \left( -\frac{(p_{x,i} - u^{(2)} m_i)^2 + p_{r,i}^2}{2kT^{(2)} m_i} \right).
\]

The function \(f_i(t > 0, -L_1 < x < L_2, p_{x,i}, p_{r,i})\) is searched by solving the Boltzmann equation. After the solution is found at the velocity grid \(S_0 = \{p_{x,\gamma}, p_{r,\gamma}\}\), gas parameters are computed as the sums given at the end of the previous section.

## 4 Presentation of macroscopic gas parameters

For the presentation of macroscopic gas parameters we use two forms: normalized form and reduced form.

Parameters presented in the first form are normalized by their values ahead of the shock wave: \(n_i / n^{(1)}_i\), \(T_i / T^{(1)}\) and so on. In the second form computed results are presented by the following reduced parameters (with the asterisks further removed):

\[
n_i^* = (n_i - n_i^{(1)}) / (n_i^{(2)} - n_i^{(1)}),
\]

\[
T_i^* = (T_i - T_i^{(1)}) / (T_i^{(2)} - T_i^{(1)}),
\]

\[
u_i^* = (u_i - u_i^{(1)}) / (u_i^{(1)} - u_i^{(2)})
\]

and similarly for the parameters of the mixture \(n, u, T\).

Characteristic parameters are the free path and velocity:

\[
l^{(1)} = \left( \frac{2\pi n^{(1)} d_1^3}{\gamma} \right)^{-1}, \quad c_0 = \sqrt{\frac{kT^{(1)}}{m_1}},
\]

\[
\tau = l^{(1)} / c_0, \quad p_0 = m_1 c_0,
\]

where \(l^{(1)}\) is the mean free path of molecules of the first component in an equilibrium state at rest with the number density \(n^{(1)}\).

## 5 Numerical results

We consider the shock wave structure in a mixture of monatomic gases assuming the hard-sphere model of molecules. The shock wave structure in the mixture is defined by a number of dimensionless parameters: Mach number \(M\), concentrations \(\chi_i^{(1)} = (n_i^{(1)}) / n^{(1)}\) of gas components before the shock wave, molecular mass ratio \(m_i / m_1\), and ratio of molecular diameters \(d_i / d_1\); \(m_1\) and \(d_1\) are parameters of the first component, \(n^{(1)}\) is the density of the mixture before the shock wave.

The ratio of diameters plays relatively small role because real molecular diameters are close to each other. The mass ratio and concentrations of components affect relaxation processes inside the shock wave and form its structure. The presentation of macroscopic values is given in the two forms: normalized form and reduced form. We use the same notation for the variables in two forms: \(n_i\) and \(n\) for number densities, \(u_i\) and \(u\) for flow velocities, \(T_i\) and \(T\) for temperatures, \(T_{xx, i}\) and \(T_{rr, i}\) for parallel
and transversal temperatures, respectively. Captions to the figures correspond to curves listed from top left to top right of each figure. In figures 1-7 we present results of calculations for a binary mixture with \( d_i/d_1 = 1 \). A heavy component is considered as the first one.

In figures Fig. 1 and Fig. 2 one can see that the differences between the profiles of densities and temperatures of components increase when the mass ratio decreases. For both cases temperature curves of the heavy component have steeper slopes than that of the light component. The temperature \( T_1 \) of the heavy component rises more quickly than the temperature of the light component \( T_2 \) and exceeds it at some point inside the shock wave. Then \( T_1 \) either approaches downstream equilibrium temperature monotonously or becomes higher than the downstream temperature and then decreases. Monotonous behavior is seen in Fig. 1, 2, and 3 where the influence of concentrations is shown. Fig. 4 shows the non-monotonous behavior of the temperature which becomes apparent at low concentrations of the heavy component and Mach number not too small. This phenomenon had already been discovered by computations in early studies [20, 21] and is known as a temperature overshoot [22, 23].

Fig. 5 shows the components of temperature tensors for the two constituents of the mixture. The
higher hump at the parallel temperature graph of the heavy gas can be explained by inertia of heavy molecules that penetrate more easily into the depth of the shock wave layer when the collisions with light molecules prevail. A contribution of the parallel component with the big hump in $T_1$ yields the overshoot of this temperature. A comparison of curves in Fig. 4 and Fig. 6 shows that the temperature overshoot increases with the rise of concentration of the light gas.

Fig. 7 presents the shock wave structure for the low mass ratio $m_2/m_1 = 0.1$. One can see a big difference of densities and temperatures of components. The temperature profile of the heavy component is much steeper than that of the light one.

Fig. 8 presents a comparison of our results for numerical density, flow velocity and temperature with computations by a different discrete ordinate method in [11]. Results of [11] are denoted by squares while our results are shown by solid curves. One can see a good agreement between both of the methods.

In [11] the computing time for one iteration step in a parallel computation, using ten CPUs on Fujitsu VPP800 computer, is 142s for $M = 2$ and 99s for $M = 3$. The computer memory for $M = 2$ is 1.7GB and for $M = 3$ is 1.4GB. In this work computations were made on a personal computer with processor Pentium 4 with the frequency 2.53GHz and
the memory 512Mb. The computing time for one iteration step for $M = 2$ is 14s and for $M = 3$ is 5s. The computer memory is 8.5Mb. For example, for Mach = 3, $m_2/m_1 = 0.5, d_2/d_1 = 1$ and various concentrations we take the following values of parameters: 11858 nodes of the momentum grid with the step $h = 0.1$, 90 nodes of the $x$-grid with the step $h_x = 0.2$, 198000 integration nodes and $\Delta t = 0.01$. With this method we can obtain the results on rough grids (1800 nodes of the momentum grid with the step $h = 0.26$, 90 nodes of the $x$-grid with the step $h_x = 0.2$, 66000 integration nodes and $\Delta t = 0.01$) with the computing time for one iteration step 1.8s.

The details of the present calculations and analysis of their accuracy can be found in our papers \[14\]. Calculations for big Mach numbers require large intervals for cylindrical coordinates in momentum space and large intervals in configuration space.

The accuracy of calculations was estimated by comparing macroscopic quantities for different grids and different numbers of integration nodes. Let $\sigma(M, S)$ represent either $n$ or $U$ or $T$ that are obtained using the grids $M$ and $S$, where $M$ is the momentum grid and $S$ is the grid in configuration space. We introduce the maximum difference between two results for two different grids $M, S$ and $M', S'$ using the formula

$$D(M', S', M, S) = \max_{\sigma = n, U, T} \left( \max_x \left| \frac{\sigma(M', S') - \sigma(M, S)}{\sigma(M, S)} \right| \right)$$

An analogous comparison was carried out for various numbers of integration nodes and fixed $M$ and $S$. The accuracy of computations is $D = 10^{-2} - 10^{-3}$. Details of our calculations with the tables containing a) data for the number density, flow velocity and temperature for the mixture with Mach = 2, 3 and concentrations $\chi_2^{(1)} = 0.1, 0.5, 0.9$; (b) grids in momentum and configuration spaces and integration nodes, together with the analysis of their accuracy can be found in our paper [14].

Fig. 9 and Fig. 10 show a comparison of our results for parallel ($T_{xx,i}$) and transversal ($T_{rr,i}$) temperatures for real gases Argon (gas 1) and Helium (gas 2) with experiments [19]. The results of [19] are marked by squares for $T_{xx,i}$ and by crosses for $T_{rr,i}$ while our
Figure 9: Parallel and transversal temperatures for Argon (gas 1) at \( M = 1.58, \frac{m_2}{m_1} = 0.1, \frac{d_2}{d_1} = 0.593, \chi_2^{(1)} = 0.9 \).

Figure 10: Parallel and transversal temperatures for Helium (gas 2) at \( M = 1.58, \frac{m_2}{m_1} = 0.1, \frac{d_2}{d_1} = 0.593, \chi_2^{(1)} = 0.9 \).

Figure 11: Temperature tensor for Argon (gas 1), Nitrogen (gas 2), Methane (gas 3), Helium (gas 4) results are denoted by solid lines. This shows a good agreement of our results with experimental data.

Fig. 11 and Fig. 12 present computational results for the three component mixture of monatomic gases: Argon (gas 1), Neon (gas 2) and Helium (gas 3), with real masses and molecular diameters. Computations were made for the following parameters of the shock wave components: \( M = 3, \frac{m_2}{m_1} = 0.1, \frac{m_3}{m_1} = 0.1, \frac{d_2}{d_1} = 0.7, \frac{d_3}{d_1} = 0.6, \chi_1^{(1)} = 0.2, \chi_2^{(1)} = 0.3, \chi_3^{(1)} = 0.5 \). On Fig. 11 one can see the maximum hump of the parallel temperature of Argon and, accordingly, on Fig. 12 the overshoot of the total temperature of Argon (the same phenomenon can be seen in Fig. 5 and Fig. 6 for a binary gas mixture) together with a big difference between the graphs of all the mixture components.

In [16] 3-component mixture has the following parameters: \( M = 3, \frac{m_1}{m_2} : \frac{m_2}{m_3} = 1 : 0.9 : 0.8, \frac{d_1}{d_2} : \frac{d_3}{d_2} = 1 : 1 : 1, \chi_1 : \chi_2 : \chi_1 = 1 : 2 : 3 \). In figures presented in this paper one can see (in our notation) the coincidence of \( n_i \) and \( n_i \), \( u_i \) and \( u_i \), \( T_{rr,i} \) for all \( i = 1, 2, 3 \) and a little difference between parallel temperatures of components \( T_{xx,i} \).

Fig. 13 and Fig. 14 show the results of computa-
Figure 12: Shock wave structure in the mixture of real gases: Argon (gas 1), Neon (gas 2), and Helium (gas 3).

Figure 13: Velocity and density for Argon (gas 1), Nitrogen (gas 2), Methane (gas 3), and Helium (gas 4) and for the mixture as a whole.

Figure 14: Shock wave structure in the mixture of real gases: Argon (gas 1), Nitrogen (gas 2), Methane (gas 3), and Helium (gas 4).

Molecular masses and diameters of the components are taken real but the internal energies of Nitrogen and Methane are not taken into account. Computations are made for \( M = 3 \) and the following parameters of the mixture:

\[
\frac{m_2}{m_1} = 0.7, \quad \frac{m_3}{m_1} = 0.4, \quad \frac{m_4}{m_1} = 0.1, \\
d_2/d_1 = 1.034, \quad d_3/d_1 = 1.144, \quad d_4/d_1 = 0.6, \\
\chi^{(1)}_1 = 0.1, \quad \chi^{(1)}_2 = 0.2, \quad \chi^{(1)}_3 = 0.3, \quad \chi^{(1)}_4 = 0.4.
\]

In Fig. 13 one can see that the graphs of the total density and velocity of the mixture lie between profiles of the components. In Fig. 14 one can see the overshoot of total temperature of Argon (gas 1) and a big difference between the graphs of all the mixture components.

Since on Fig. 12 and Fig. 14 the ratios of molecular masses for the last component differ too much \((m_i/m_1 = 0.1, \ i = 3, 4)\), we see that the profile of temperature of the last (lightest) component is flat. If the mass ratio of the last component differs not so much from the others, we do not see so flat profiles. All the calculations were performed on the laptop computer Sony VAIO, processor Intel(R) Core(TM)2CPU, 1.66GHz+1.66GHz, 1.00GB.
of RAM. For these calculations we have taken the following values of parameters: 20000 nodes of the momentum grid with the step $h = 0.08$, 200 nodes of the $x$-grid with the step $h_x = 0.15$, 66000 integration nodes and $\Delta t = 0.01$.

Thus, we have made calculations for two, three and four components of the mixture. The test calculations were made for binary gas mixtures with the concentration $\chi^{(2)}_2$ of the second component: 0.1, 0.5, 0.9, 0.95, ratio of masses 0.1, 0.25, 0.5, Mach number 1.5, 2, 3, 6, 8.

The method of [8] was extended for the gases with internal degrees of freedom in [24], where it can incorporate real physical parameters of molecular potentials and internal energy spectrum.

6 Conclusions

Computational results for the shock wave structure presented in this paper were obtained by a unique approach based on the application of the Conservative Projection Method (CPM) [7][10] for solving the classical kinetic Boltzmann equation for monatomic gases. This method was extended to binary gas mixtures in cylindrical coordinates and, later on, to three and four component gas mixtures [1,14,15] and in this paper. This method ensures strict conservation of mass, momentum and energy. The transition from the upstream to downstream state was presented by distribution functions and their moments (macroscopic values) for various parameters (Mach numbers, ratios of masses and concentrations). Details of our calculations together with analysis of the mixture accuracy can be found in paper [14]. With this method we can obtain results on rough grids which coincide well with the results obtained on more fine grids. The numerical results have been compared with numerical and experimental results of other authors with a good agreement with them. All computations were performed on personal computers without using parallel processing. We have shown that the projection method for a gas mixture solves the shock wave problem with acceptable precision, small time of calculations and small computer memory.

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