# Physically Based Modeling Rigid Body Simulation

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# **Rigid Body Simulation**

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## Introduction

This portion of the course notes deals with the problem of rigid body dynamics. To help get you started simulating rigid body motion, we've provided code fragments that implement most of the concepts discussed in these notes. This segment of the course notes is divided into two parts. The first part covers the motion of rigid bodies that are completely *unconstrained* in their allowable motion; that is, simulations that aren't concerned about collisions between rigid bodies. Given any external forces acting on a rigid body, we'll show how to simulate the motion of the body in response to these forces. The mathematical derivations in these notes are meant to be fairly informal and intuitive.

The second part of the notes tackles the problem of *constrained* motion that arises when we regard bodies as solid, and need to disallow inter-penetration. We enforce these non-penetration constraints by computing appropriate contact forces between contacting bodies. Given values for these contact forces, simulation proceeds exactly as in the unconstrained case: we simply apply all the forces to the bodies and let the simulation unfold as though the motions of bodies are completely unconstrained. If we have computed the contact forces correctly, the resulting motion of the bodies will be free from inter-penetration. The computation of these contact forces is the most demanding component of the entire simulation process.<sup>1</sup>

<sup>&</sup>lt;sup>1</sup>Collision detection (i.e. determining the points of contact between bodies) runs a close second though!

# Part I. Unconstrained Rigid Body Dynamics

#### 1 Simulation Basics

This portion of the course notes is geared towards a full implementation of rigid body motion. In this section, we'll show the basic structure for simulating the motion of a rigid body. In section 2, we'll define the terms, concepts, and equations we need to implement a rigid body simulator. Following this, we'll give some code to actually implement the equations we need. Derivations for some of the concepts and equations we will be using will be left to appendix A.

The only thing you need to be familiar with at this point are the basic concepts (but not the numerical details) of solving ordinary differential equations. If you're not familiar with this topic, you're in luck: just turn back to the beginning of these course notes, and read the section on "Differential Equation Basics." You also might want to read the next section on "Particle Dynamics" as well, although we're about to repeat some of that material here anyway.

Simulating the motion of a rigid body is almost the same as simulating the motion of a particle, so let's start with particle simulation. The way we simulate a particle is as follows. We let a function x(t) denote the particle's location in world space (the space all particles or bodies occupy during simulation) at time t. The function  $v(t) = \dot{x}(t) = \frac{d}{dt}x(t)$  gives the velocity of the particle at time t. The state of a particle at time t is the particle's position and velocity. We generalize this concept by defining a state vector  $\mathbf{X}(t)$  for a system: for a single particle,

$$\mathbf{X}(t) = \begin{pmatrix} x(t) \\ v(t) \end{pmatrix}. \tag{1-1}$$

When we're talking about an actual implementation, we have to "flatten" out  $\mathbf{X}(t)$  into an array. For a single particle,  $\mathbf{X}(t)$  can be described as an array of six numbers: typically, we'd let the first three elements of the array represent x(t), and the last three elements represent v(t). Later, when we talk about state vectors  $\mathbf{X}(t)$  that contain matrices as well as vectors, the same sort of operation is done to flatten  $\mathbf{X}(t)$  into an array. Of course, we'll also have to reverse this process and turn an array of numbers back into a state vector  $\mathbf{X}(t)$ . This all comes down to pretty simple bookkeeping though, so henceforth, we'll assume that we know how to convert any sort of state vector  $\mathbf{X}(t)$  to an array (of the appropriate length) and vice versa. (For a simple example involving particles, look through the "Particle System Dynamics" section of these notes.)

For a system with n particles, we enlarge  $\mathbf{X}(t)$  to be

$$\mathbf{X}(t) = \begin{pmatrix} x_1(t) \\ v_1(t) \\ \vdots \\ x_n(t) \\ v_n(t) \end{pmatrix}$$
 (1-2)

where  $x_i(t)$  and  $v_i(t)$  are the position and velocity of the *i*th particle. Working with *n* particles is no harder than working with one particle, so we'll let  $\mathbf{X}(t)$  be the state vector for a single particle for now (and when we get to it later, a single rigid body).

To actually simulate the motion of our particle, we need to know one more thing—the force acting on the particle at time t. We'll define F(t) as the force acting on our particle at time t. The function F(t) is the sum of all the forces acting on the particle: gravity, wind, spring forces, etc. If the particle has mass m, then the change of  $\mathbf{X}$  over time is given by

$$\frac{d}{dt}\mathbf{X}(t) = \frac{d}{dt} \begin{pmatrix} x(t) \\ v(t) \end{pmatrix} = \begin{pmatrix} v(t) \\ F(t)/m \end{pmatrix}. \tag{1-3}$$

Given any value of  $\mathbf{X}(t)$ , equation (1–3) describes how  $\mathbf{X}(t)$  is instantaneously changing at time t. A simulation starts with some initial conditions for  $\mathbf{X}(0)$ , (i.e. values for x(0) and v(0)) and then uses a numerical equation solver to track the change or "flow" of  $\mathbf{X}$  over time, for as long as we're interested in. If all we want to know is the particle's location one second from now, we ask the solver to compute  $\mathbf{X}(1)$ , assuming that time units are in seconds. If we're going to animate the motion of the particle, we'd want to compute  $\mathbf{X}(\frac{1}{30})$ ,  $\mathbf{X}(\frac{2}{30})$  and so on.

The numerical method used by the solver is relatively unimportant with respect to our actual implementation. Let's look at how we'd actually interact with a numerical solver, in a C++-like language. Assume we have access to a numerical solver, which we'll generically write as a function named ode. Typically, ode has the following specification:

We pass an initial state vector to ode as an array  $\times 0$ . The solver ode knows nothing about the inherent structure of  $\times 0$ . Since solvers can handle problems of arbitrary dimension, we also have to pass the length len of  $\times 0$ . (For a system of n particles, we'd obviously have len = 6n.) We also pass the solver the starting and ending times of the simulation, t0 and t1. The solver's goal is to compute the state vector at time t1 and return it in the array  $\times End$ .

We also pass a function Dxdt() to ode. Given an array y that encodes a state vector  $\mathbf{X}(t)$  and a time t, Dxdt() must compute and return  $\frac{d}{dt}\mathbf{X}(t)$  in the array xdot. (The reason we must pass t to Dxdt() is that we may have time-varying forces acting in our system. In that case, Dxdt() would have to know "what time it is" to determine the value of those forces.) In tracing the flow of  $\mathbf{X}(t)$  from t0 to t1, the solver ode is allowed to call Dxdt() as often as it likes. Given that we have such a routine ode, the only work we need to do is to code up the routine Dxdt() which we'll give as a parameter to ode.

Simulating rigid bodies follows exactly the same mold as simulating particles. The only difference is that the state vector  $\mathbf{X}(t)$  for a rigid body holds more information, and the derivative  $\frac{d}{dt}\mathbf{X}(t)$  is a little more complicated. However, we'll use exactly the same paradigm of tracking the movement of a rigid body using a solver ode, which we'll supply with a function Dxdt().

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# 2 Rigid Body Concepts

The goal of this section is to develop an analogue to equation (1-3), for rigid bodies. The final differential equation we develop is given in section 2.11. In order to do this though, we need to define a lot of concepts first and relations first. Some of the longer derivations are found in appendix A. In the next section, we'll show how to write the function Dxdt() needed by the numerical solver ode to compute the derivative  $\frac{d}{dt}X(t)$  developed in this section.

#### 2.1 Position and Orientation

The location of a particle in space at time t can be described as a vector x(t), which describes the translation of the particle from the origin. Rigid bodies are more complicated, in that in addition to translating them, we can also rotate them. To locate a rigid body in world space, we'll use a vector x(t), which describes the translation of the body. We must also describe the rotation of the body, which we'll do (for now) in terms of a  $3 \times 3$  rotation matrix R(t). We will call x(t) and R(t) the spatial variables of a rigid body.

A rigid body, unlike a particle, occupies a volume of space and has a particular shape. Because a rigid body can undergo only rotation and translation, we define the shape of a rigid body in terms of a fixed and unchanging space called *body space*. Given a geometric description of the body in body space, we use x(t) and R(t) to transform the body-space description into world space (figure 1). In order to simplify some equations we'll be using, we'll require that our description of the rigid body in body space be such that the *center of mass* of the body lies at the origin, (0,0,0). We'll define the center of mass more precisely later, but for now, the center of mass can be thought of as a point in the rigid body that lies at the geometric center of the body. In describing the body's shape, we require that this geometric center lie at (0,0,0) in body space. If we agree that R(t) specifies a rotation of the body about the center of mass, then a fixed vector r in body space will be rotated to the world-space vector R(t)r at time t. Likewise, if  $p_0$  is an arbitrary point on the rigid body, in body space, then the world-space location p(t) of  $p_0$  is the result of first rotating  $p_0$  about the origin and then translating it:

$$p(t) = R(t)p_0 + x(t). (2-1)$$

Since the center of mass of the body lies at the origin, the world-space location of the center of mass is always given directly by x(t). This lets us attach a very physical meaning to x(t) by saying that x(t) is the location of the center of mass in world space at time t. We can also attach a physical meaning to R(t). Consider the x axis in body space i.e. the vector (1,0,0). At time t, this vector has direction

$$R(t) \left( \begin{array}{c} 1 \\ 0 \\ 0 \end{array} \right)$$

in world space. If we write out the components of R(t) as

$$R(t) = \begin{pmatrix} r_{xx} & r_{yx} & r_{zx} \\ r_{xy} & r_{yy} & r_{zy} \\ r_{xz} & r_{yz} & r_{zz} \end{pmatrix},$$
(2-2)

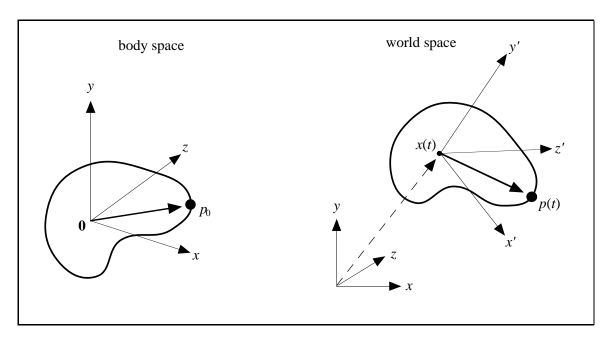


Figure 1: The center of mass is transformed to the point x(t) in world space, at time t. The fixed x, y, and z axes of the body in body space transform to the vectors x' = R(t)x, y' = R(t)y and z' = R(t)z. The fixed point  $p_0$  in body space is transformed to the point  $p(t) = R(t)p_0 + x(t)$ .

then

$$R(t) \begin{pmatrix} 1 \\ 0 \\ 0 \end{pmatrix} = \begin{pmatrix} r_{xx} \\ r_{xy} \\ r_{xz} \end{pmatrix}$$
 (2-3)

which is the first column of R(t). The physical meaning of R(t) is that R(t)'s first column gives the direction that the rigid body's x axis points in, when transformed to world space at time t. Similarly, the second and third columns of R(t),

$$\left( \begin{array}{c} r_{yx} \\ r_{yy} \\ r_{yz} \end{array} \right)$$
 and  $\left( \begin{array}{c} r_{zx} \\ r_{zy} \\ r_{zz} \end{array} \right)$ 

are the directions of the y and z axes of the rigid body in world space at time t (figure 2).

#### 2.2 Linear Velocity

For simplicity, we'll call x(t) and R(t) the *position* and *orientation* of the body at time t. The next thing we need to do is define how the position and orientation change over time. This means we need expressions for  $\dot{x}(t)$  and  $\dot{R}(t)$ . Since x(t) is the position of the center of mass in world space,  $\dot{x}(t)$  is the velocity of the center of mass in world space. We'll define the *linear velocity* v(t) as this velocity:

$$v(t) = \dot{x}(t). \tag{2-4}$$

If we imagine that the orientation of the body is fixed, then the only movement the body can undergo is a pure translation. The quantity v(t) gives the velocity of this translation.

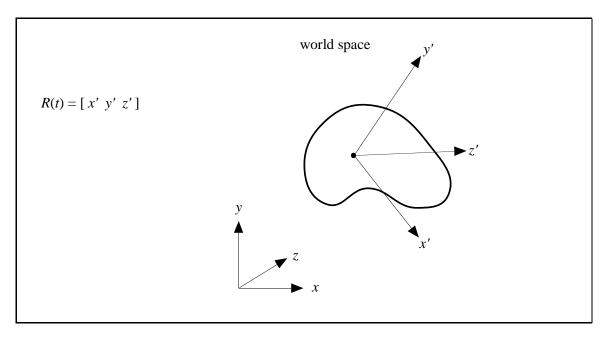


Figure 2: Physical interpretation of the orientation matrix R(t). At time t, the columns of R(t) are the world-space directions that the body-space x, y, and z axes transform to.

#### 2.3 Angular Velocity

In addition to translating, a rigid body can also spin. Imagine however that we freeze the position of the center of mass in space. Any movement of the points of the body must therefore be due to the body spinning about some axis that passes through the center of mass. (Otherwise the center of mass would itself be moving). We can describe that spin as a vector  $\omega(t)$ . The *direction* of  $\omega(t)$  gives the direction of the axis about which the body is spinning (figure 3). The magnitude of  $\omega(t)$ ,  $|\omega(t)|$ , tells how fast the body is spinning.  $|\omega(t)|$  has the dimensions of revolutions/time; thus,  $|\omega(t)|$  relates the angle through which the body will rotate over a given period of time, if the angular velocity remains constant. The quantity  $\omega(t)$  is called the *angular velocity*.

For linear velocity, x(t) and v(t) are related by  $v(t) = \frac{d}{dt}x(t)$ . How are R(t) and  $\omega(t)$  related? (Clearly,  $\dot{R}(t)$  cannot be  $\omega(t)$ , since R(t) is a matrix, and  $\omega(t)$  is a vector.) To answer this question, let's remind ourselves of the physical meaning of R(t). We know that the columns of R(t) tell us the directions of the transformed x, y and z body axes at time t. That means that the columns of  $\dot{R}(t)$  must describe the *velocity* with which the x, y, and z axes are being transformed. To discover the relationship between  $\omega(t)$  and R(t), let's examine how the change of an arbitrary vector in a rigid body is related to the angular velocity  $\omega(t)$ .

Figure 4 shows a rigid body with angular velocity  $\omega(t)$ . Consider a vector r(t) at time t specified in world space. Suppose that we consider this vector fixed to the body; that is, r(t) moves along with the rigid body through world space. Since r(t) is a direction, it is independent of any translational effects; in particular,  $\dot{r}(t)$  is independent of v(t). To study  $\dot{r}(t)$ , we decompose r(t) into vectors a and b, where a is parallel to  $\omega(t)$  and b is perpendicular to  $\omega(t)$ . Suppose the rigid body were to maintain a constant angular velocity, so that the tip of r(t) traces out a circle centered on the  $\omega(t)$  axis (figure 4). The radius of this circle is |b|. Since the tip of the vector r(t) is instantaneously moving along this circle, the instantaneous change of r(t) is perpendicular to both b and  $\omega(t)$ . Since the tip of r(t) is moving in a circle of radius b, the instantaneous velocity of r(t) has magnitude  $|b||\omega(t)|$ .

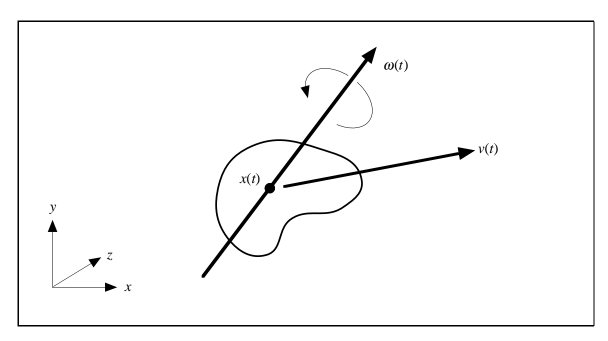


Figure 3: Linear velocity v(t) and angular velocity  $\omega(t)$  of a rigid body.

Since b and  $\omega(t)$  are perpendicular, their cross product has magnitude

$$|\omega(t) \times b| = |\omega(t)| |b|. \tag{2-5}$$

Putting this together, we can write  $\dot{r}(t) = \omega(t) \times (b)$ . However, since r(t) = a + b and a is parallel to  $\omega(t)$ , we have  $\omega(t) \times a = 0$  and thus

$$\dot{r}(t) = \omega(t) \times b = \omega(t) \times b + \omega(t) \times a = \omega(t) \times (b+a). \tag{2-6}$$

Thus, we can simply express the rate of change of a vector as

$$\dot{r}(t) = \omega(t) \times r(t). \tag{2-7}$$

Let's put all this together now. At time t, we know that the direction of the x axis of the rigid body in world space is the first column of R(t), which is

$$\begin{pmatrix} r_{xx} \\ r_{xy} \\ r_{xz} \end{pmatrix}$$
.

At time t, the derivative of the first column of R(t) is just the rate of change of this vector: using the cross product rule we just discovered, this change is

$$\omega(t) \times \left( \begin{array}{c} r_{xx} \\ r_{xy} \\ r_{xz} \end{array} \right).$$

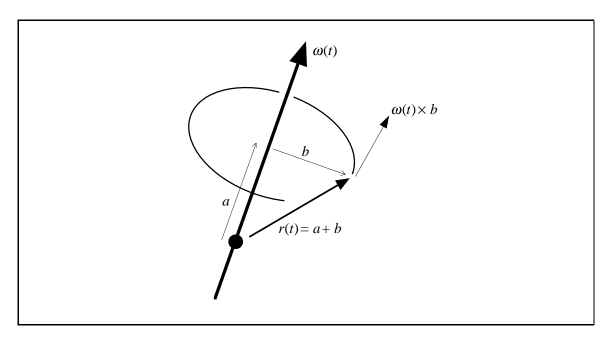


Figure 4: The rate of change of a rotating vector. As the tip of r(t) spins about the  $\omega(t)$  axis, it traces out a circle of diameter |b|. The speed of the tip of r(t) is  $|\omega(t)||b|$ .

The same obviously holds for the other two columns of R(t). This means that we can write

$$\dot{R} = \left(\omega(t) \times \begin{pmatrix} r_{xx} \\ r_{xy} \\ r_{xz} \end{pmatrix} \quad \omega(t) \times \begin{pmatrix} r_{yx} \\ r_{yy} \\ r_{yz} \end{pmatrix} \quad \omega(t) \times \begin{pmatrix} r_{zx} \\ r_{zy} \\ r_{zz} \end{pmatrix} \right). \tag{2-8}$$

This is too cumbersome an expression to tote around though. To simplify things, we'll use the following trick. If a and b are 3-vectors, then  $a \times b$  is the vector

$$\begin{pmatrix} a_y b_z - b_y a_z \\ -a_x b_z + b_x a_z \\ a_x b_y - b_x a_y \end{pmatrix}.$$

Given the vector a, let us define  $a^*$  to be the matrix

$$\left(\begin{array}{ccc}
0 & -a_z & a_y \\
a_z & 0 & -a_x \\
-a_y & a_x & 0
\end{array}\right).$$

Then<sup>2</sup>

$$a^*b = \begin{pmatrix} 0 & -a_z & a_y \\ a_z & 0 & -a_x \\ -a_y & a_x & 0 \end{pmatrix} \begin{pmatrix} b_x \\ b_y \\ b_z \end{pmatrix} = \begin{pmatrix} a_y b_z - b_y a_z \\ -a_x b_z + b_x a_z \\ a_x b_y - b_x a_y \end{pmatrix} = a \times b.$$
 (2-9)

<sup>&</sup>lt;sup>2</sup>This looks a little too "magical" at first. Did someone discover this identity accidentally? Is it a relation that just happens to work? This construct can be derived by considering what's known as *infinitesimal* rotations. The interested reader might wish to read chapter 4.8 of Goldstein[10] for a more complete derivation of the  $a^*$  matrix.

Using the "\*" notation, we can rewrite  $\dot{R}(t)$  more simply as

$$\dot{R}(t) = \left(\omega(t)^* \begin{pmatrix} r_{xx} \\ r_{xy} \\ r_{xz} \end{pmatrix} \quad \omega(t)^* \begin{pmatrix} r_{yx} \\ r_{yy} \\ r_{yz} \end{pmatrix} \quad \omega(t)^* \begin{pmatrix} r_{zx} \\ r_{zy} \\ r_{zz} \end{pmatrix} \right). \tag{2-10}$$

By the rules of matrix multiplication, we can factor this into

$$\dot{R}(t) = \omega(t)^* \begin{pmatrix} r_{xx} \\ r_{xy} \\ r_{xz} \end{pmatrix} \begin{pmatrix} r_{yx} \\ r_{yy} \\ r_{yz} \end{pmatrix} \begin{pmatrix} r_{zx} \\ r_{zy} \\ r_{zz} \end{pmatrix}$$
(2-11)

which is a matrix-matrix multiplication. But since the matrix on the right is R(t) itself, we get simply that

$$\dot{R}(t) = \omega(t)^* R(t). \tag{2-12}$$

This, at last, gives us the relation we wanted between  $\dot{R}(t)$  and  $\omega(t)$ . Note the correspondence between  $\dot{r}(t) = \omega(t) \times r(t)$  for a vector, and  $\dot{R}(t) = \omega(t)^* R(t)$  for the rotation matrix.

#### 2.4 Mass of a Body

In order to work out some derivations, we'll need to (conceptually) perform some integrations over the volume of our rigid body. To make these derivations simpler, we're going to temporarily imagine that a rigid body is made up of a large number of small particles. The particles are indexed from 1 to N. The mass of the ith particle is  $m_i$ , and each particle has a (constant) location  $r_{0i}$  in body space. The location of the ith particle in world space at time t, denoted  $r_i(t)$ , is therefore given by the formula

$$r_i(t) = R(t)r_{0i} + x(t).$$
 (2–13)

The total mass of the body, M, is the sum

$$M = \sum_{i=1}^{N} m_i. (2-14)$$

(Henceforth, summations are assumed to be summed from 1 to N with index variable i.)

#### 2.5 Velocity of a Particle

The *velocity*  $\dot{r}_i(t)$  of the *i*th particle is obtained by differentiating equation (2–13): using the relation  $\dot{R}(t) = \omega^* R(t)$ , we obtain

$$\dot{r}_i(t) = \omega^* R(t) r_{0i} + v(t). \tag{2-15}$$

We can rewrite this as

$$\dot{r}_i(t) = \omega(t)^* R(t) r_{0i} + v(t)$$

$$= \omega(t)^* (R(t) r_{0i} + x(t) - x(t)) + v(t)$$

$$= \omega(t)^* (r_i(t) - x(t)) + v(t)$$
(2-16)

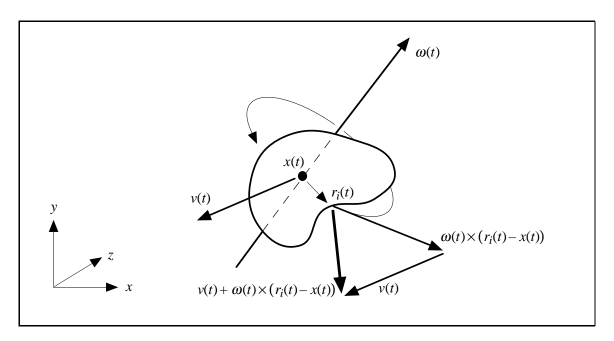


Figure 5: The velocity of the *i*th point of a rigid body in world space. The velocity of  $r_i(t)$  can be decomposed into a linear term v(t) and an angular term  $\omega(t) \times (r_i(t) - x(t))$ .

using the definition of  $r_i(t)$  from equation (2–13). Recall from the definition of the "\*" operator that  $\omega(t)^*a = \omega(t) \times a$  for any vector a. Using this, we can simply write

$$\dot{r}_i(t) = \omega(t) \times (r_i(t) - x(t)) + v(t).$$
 (2–17)

Note that this separates the velocity of a point on a rigid body into two components (figure 5): a linear component v(t), and an angular component  $\omega \times (r_i(t) - x(t))$ .

#### 2.6 Center of Mass

Our definition of the center of mass is going to enable us to likewise separate the dynamics of bodies into linear and angular components. The center of mass of a body in world space is defined to be

$$\frac{\sum m_i r_i(t)}{M} \tag{2-18}$$

where M is the mass of the body (i.e. the sum of the individual masses  $m_i$ ). When we say that we are using a center of mass coordinate system, we mean that in body space,

$$\frac{\sum m_i r_{0i}}{M} = \mathbf{0} = \begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}. \tag{2-19}$$

Note that this implies that  $\sum m_i r_{0i} = \mathbf{0}$  as well.

We have spoken of x(t) as being the location of the center of mass at time t. Is this true? Yes: since the ith particle has position  $r_i(t) = R(t)r_{0i} + x(t)$  at time t, the center of mass at time t is

$$\frac{\sum m_i r_i(t)}{M} = \frac{\sum m_i (R(t) r_{0i} + x(t))}{M} = \frac{R(t) \sum m_i r_{0i} + \sum m_i x(t)}{M} = x(t) \frac{\sum m_i}{M} = x(t).$$

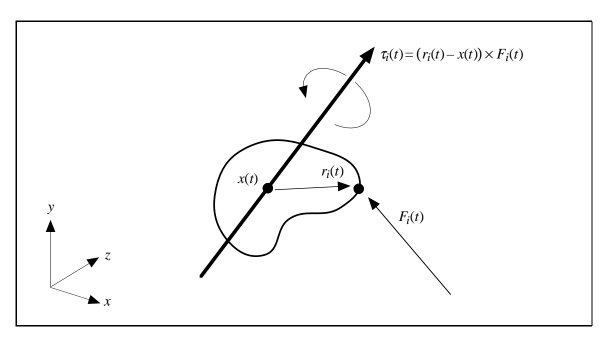


Figure 6: The torque  $\tau_i(t)$  due to a force  $F_i(t)$  acting at  $r_i(t)$  on a rigid body.

Additionally, the relation

$$\sum m_i(r_i(t) - x(t)) = \sum m_i(R(t)r_{0i} + x(t) - x(t)) = R(t) \sum m_i r_{0i} = \mathbf{0}$$
 (2-20)

is also very useful.

## 2.7 Force and Torque

When we imagine a force acting on a rigid body due to some external influence (e.g. gravity, wind, contact forces), we imagine that the force acts on a particular particle of the body. (Remember that our particle model is conceptual only. We can have a force act at any geometrical location on or inside the body, because we can always imagine that there happens to be a particle at that exact location.) The location of the particle the force acts on defines the location at which the force acts. We will let  $F_i(t)$  denote the total force from external forces acting on the *i*th particle at time *t*. Also, we define the external *torque*  $\tau_i(t)$  acting on the *i*th particle as

$$\tau_i(t) = (r_i(t) - x(t)) \times F_i(t).$$
 (2–21)

Torque differs from force in that the torque on a particle depends on the location  $r_i(t)$  of the particle, relative to the center of mass x(t). We can intuitively think of the direction of  $\tau_i(t)$  as being the axis the body would spin about due to  $F_i(t)$ , if the center of mass were held firmly in place (figure 6).

The total external force F(t) acting on the body is the sum of the  $F_i(t)$ :

$$F(t) = \sum F_i(t) \tag{2-22}$$

while the total external torque is defined similarly as

$$\tau(t) = \sum \tau_i(t) = \sum (r_i(t) - x(t)) \times F_i(t).$$
 (2-23)

Note that F(t) conveys no information about where the various forces acted on the body; however,  $\tau(t)$  does tell us something about the distribution of the forces  $F_i(t)$  over the body.

#### 2.8 Linear Momentum

The linear momentum p of a particle with mass m and velocity v is defined as

$$p = mv. (2-24)$$

The total linear momentum P(t) of a rigid body is the sum of the products of the mass and velocity of each particle:

$$P(t) = \sum m_i \dot{r}_i(t). \tag{2-25}$$

From equation (2–17), the velocity  $\dot{r}_i(t)$  of the *i*th particle is  $\dot{r}_i(t) = v(t) + \omega(t) \times (r_i(t) - x(t))$ . Thus, the total linear momentum of the body is

$$P(t) = \sum m_i \dot{r}_i(t)$$

$$= \sum \left( m_i v(t) + m_i \omega(t) \times (r_i(t) - x(t)) \right)$$

$$= \sum m_i v(t) + \omega(t) \times \sum m_i (r_i(t) - x(t)).$$
(2-26)

Because we are using a center of mass coordinate system, we can apply equation (2–20) and obtain

$$P(t) = \sum m_i v(t) = \left(\sum m_i\right) v(t) = Mv(t). \tag{2-27}$$

This gives us the nice result that the total linear momentum of our rigid body is the same as if the body was simply a particle with mass M and velocity v(t). Because of this, we have a simple transformation between P(t) and v(t): P(t) = Mv(t) and v(t) = P(t)/M. Since M is a constant,

$$\dot{v}(t) = \frac{\dot{P}(t)}{M}.\tag{2-28}$$

The concept of linear momentum lets us express the effect of the total force F(t) on a rigid body quite simply. Appendix A derives the relation

$$\dot{P}(t) = F(t) \tag{2-29}$$

which says that the change in linear momentum is equivalent to the total force acting on a body. Note that P(t) tells us nothing about the rotational velocity of a body, which is good, because F(t) also conveys nothing about the change of rotational velocity of a body!

Since the relationship between P(t) and v(t) is simple, we will be using P(t) as a state variable for our rigid body, instead of v(t). We could of course let v(t) be a state variable, and use the relation

$$\dot{v}(t) = \frac{F(t)}{M}. (2-30)$$

However, using P(t) instead of v(t) as a state variable will be more consistent with the way we will be dealing with angular velocity and acceleration.

#### 2.9 Angular Momentum

While the concept of linear momentum is pretty intuitive (P(t) = Mv(t)), the concept of angular momentum (for a rigid body) is not. The only reason that one even bothers with the angular momentum of a rigid body is that it lets you write simpler equations than you would get if you stuck with angular velocity. With that in mind, it's probably best not to worry about attaching an intuitive physical explanation to angular momentum—all in all, it's a most unintuitive concept. Angular momentum ends up simplifying equations because it is conserved in nature, while angular *velocity* is not: if you have a body floating through space with no torque acting on it, the body's angular momentum is constant. This is not true for a body's angular velocity though: even if the angular momentum of a body is constant, the body's angular *velocity* may not be! Consequently, a body's angular velocity can vary even when no force acts on the body. Because of this, it ends up being simpler to choose angular momentum as a state variable over angular velocity.

For linear momentum, we have the relation P(t) = Mv(t). Similarly, we define the total angular momentum L(t) of a rigid body by the equation  $L(t) = I(t)\omega(t)$ , where I(t) is a  $3 \times 3$  matrix (technically a rank-two tensor) called the *inertia tensor*, which we will describe momentarily. The inertia tensor I(t) describes how the mass in a body is distributed relative to the body's center of mass. The tensor I(t) depends on the orientation of a body, but does not depend on the body's translation. Note that for both the angular and the linear case, momentum is a linear function of velocity—it's just that in the angular case the scaling factor is a matrix, while it's simply a scalar in the linear case. Note also that L(t) is independent of any translational effects, while P(t) is independent of any rotational effects.

The relationship between L(t) and the total torque  $\tau(t)$  is very simple: appendix A derives

$$\dot{L}(t) = \tau(t), \tag{2-31}$$

analogous to the relation  $\dot{P}(t) = F(t)$ .

#### 2.10 The Inertia Tensor

The inertia tensor I(t) is the scaling factor between angular momentum L(t) and angular velocity  $\omega(t)$ . At a given time t, let  $r'_i$  be the displacement of the ith particle from x(t) by defining  $r'_i = r_i(t) - x(t)$ . The tensor I(t) is expressed in terms of  $r'_i$  as the symmetric matrix

$$I(t) = \sum \begin{pmatrix} m_i (r'_{iy}^2 + r'_{iz}^2) & -m_i r'_{ix} r'_{iy} & -m_i r'_{ix} r'_{iz} \\ -m_i r'_{iy} r'_{ix} & m_i (r'_{ix}^2 + r'_{iz}^2) & -m_i r'_{iy} r'_{iz} \\ -m_i r'_{iz} r'_{ix} & -m_i r'_{iz} r'_{iy} & m_i (r'_{ix}^2 + r'_{iy}^2) \end{pmatrix}$$
(2-32)

For an actual implementation, we replace the finite sums with integrals over a body's volume in world space. The mass terms  $m_i$  are replaced by a density function. At first glance, it seems that we would need to evaluate these integrals to find I(t) whenever the orientation R(t) changes. This would be prohibitively expensive to do during a simulation unless the body's shape was so simple (for example, a sphere or cube) that that the integrals could be evaluated symbolically.

Fortunately, by using body-space coordinates we can cheaply compute the inertia tensor for any orientation R(t) in terms of a precomputed integral in body-space coordinates. (This integral is typically computed before the simulation begins and should be regarded as one of the input parameters

describing a physical property of the body.) Using the fact that  $r_i^T r_i' = r_{ix}'^2 + r_{iy}'^2 + r_{iz}'^2$ , we can rewrite I(t) as the difference

$$I(t) = \sum_{i} m_{i} r_{i}^{\prime T} r_{i}^{\prime} \begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix} - \begin{pmatrix} m_{i} r_{ix}^{\prime 2} & m_{i} r_{ix}^{\prime} r_{iy}^{\prime} & m_{i} r_{ix}^{\prime} r_{iz}^{\prime} \\ m_{i} r_{iy}^{\prime} r_{ix}^{\prime} & m_{i} r_{iy}^{\prime 2} & m_{i} r_{iy}^{\prime} r_{iz}^{\prime} \\ m_{i} r_{iz}^{\prime} r_{ix}^{\prime} & m_{i} r_{iz}^{\prime} r_{iy}^{\prime} & m_{i} r_{iz}^{\prime 2} \end{pmatrix}$$
(2-33)

Taking the outer product multiplication of  $r'_i$  with itself, that is

$$r_{i}'r_{i}'^{T} = \begin{pmatrix} r_{ix}' \\ r_{iy}' \\ r_{iz}' \end{pmatrix} \begin{pmatrix} r_{ix}' & r_{iy}' & r_{iz}' \\ r_{iy}' & r_{iz}' \end{pmatrix} = \begin{pmatrix} r_{ix}'^{2} & r_{ix}'r_{iy}' & r_{ix}'r_{iz}' \\ r_{iy}'r_{ix}' & r_{iz}'^{2} & r_{ix}'r_{iz}' \\ r_{iz}'r_{ix}' & r_{iz}'r_{iy}' & r_{iz}'^{2} \end{pmatrix}$$
(2-34)

and letting 1 denote the  $3 \times 3$  identity matrix, we can express I(t) simply as

$$I(t) = \sum m_i((r_i^{'T}r_i^{'})\mathbf{1} - r_i^{'}r_i^{'T})$$
 (2-35)

How does this help?

Since  $r_i(t) = R(t)r_{0i} + x(t)$  where  $r_{0i}$  is a constant,  $r'_i = R(t)r_{0i}$ . Then, since  $R(t)R(t)^T = 1$ ,

$$I(t) = \sum_{i} m_{i} ((r_{i}^{T} r_{i}^{\prime}) \mathbf{1} - r_{i}^{\prime} r_{i}^{T})$$

$$= \sum_{i} m_{i} ((R(t) r_{0i})^{T} (R(t) r_{0i}) \mathbf{1} - (R(t) r_{0i}) (R(t) r_{0i})^{T})$$

$$= \sum_{i} m_{i} (r_{0i}^{T} R(t)^{T} R(t) r_{0i} \mathbf{1} - R(t) r_{0i} r_{0i}^{T} R(t)^{T})$$

$$= \sum_{i} m_{i} ((r_{0i}^{T} r_{0i}) \mathbf{1} - R(t) r_{0i} r_{0i}^{T} R(t)^{T}).$$
(2-36)

Since  $r_0^T r_{0i}$  is a scalar, we can rearrange things by writing

$$I(t) = \sum_{i} m_{i}((r_{0i}^{T} r_{0i}) \mathbf{1} - R(t) r_{0i} r_{0i}^{T} R(t)^{T})$$

$$= \sum_{i} m_{i}(R(t) (r_{0i}^{T} r_{0i}) R(t)^{T} \mathbf{1} - R(t) r_{0i} r_{0i}^{T} R(t)^{T})$$

$$= R(t) \left( \sum_{i} m_{i}((r_{0i}^{T} r_{0i}) \mathbf{1} - r_{0i} r_{0i}^{T}) \right) R(t)^{T}.$$
(2-37)

If we define  $I_{body}$  as the matrix

$$I_{body} = \sum m_i ((r_{0i}^T r_{0i}) \mathbf{1} - r_{0i} r_{0i}^T)$$
 (2-38)

then from the previous equation we have

$$I(t) = R(t)I_{body}R(t)^{T}. (2-39)$$

Since  $I_{body}$  is specified in body-space, it is constant over the simulation. Thus, by precomputing  $I_{body}$  for a body before the simulation begins, we can easily compute I(t) from  $I_{body}$  and the orientation matrix R(t). Section 5.1 derives the body-space inertia tensor for a rectangular object in terms of an integral over the body's volume in body space.

Also, the inverse of I(t) is given by the formula

$$I^{-1}(t) = (R(t)I_{body}R(t)^{T})^{-1}$$

$$= (R(t)^{T})^{-1}I_{body}^{-1}R(t)^{-1}$$

$$= R(t)I_{body}^{-1}R(t)^{T}$$
(2-40)

since, for rotation matrices,  $R(t)^T = R(t)^{-1}$  and  $\left(R(t)^T\right)^T = R(t)$ . Clearly,  $I_{body}^{-1}$  is also a constant during the simulation.

#### 2.11 Rigid Body Equations of Motion

Finally, we have covered all the concepts we need to define the state vector  $\mathbf{X}(t)$ ! For a rigid body, we will define  $\mathbf{X}(t)$  as

$$\mathbf{X}(t) = \begin{pmatrix} x(t) \\ R(t) \\ P(t) \\ L(t) \end{pmatrix}. \tag{2-41}$$

Thus, the state of a rigid body is its position and orientation (describing spatial information), and its linear and angular momentum (describing velocity information). The mass M of the body and body-space inertia tensor  $I_{body}$  are constants, which we assume we know when the simulation begins. At any given time, the auxiliary quantities I(t),  $\omega(t)$  and v(t) are computed by

$$v(t) = \frac{P(t)}{M}, \quad I(t) = R(t)I_{body}R(t)^{T}$$
 and  $\omega(t) = I(t)^{-1}L(t).$  (2-42)

The derivative  $\frac{d}{dt}\mathbf{X}(t)$  is

$$\frac{d}{dt}\mathbf{X}(t) = \frac{d}{dt} \begin{pmatrix} x(t) \\ R(t) \\ P(t) \\ L(t) \end{pmatrix} = \begin{pmatrix} v(t) \\ \omega(t)^*R(t) \\ F(t) \\ \tau(t) \end{pmatrix}. \tag{2-43}$$

The next section gives an implementation for the function Dxdt ( ) that computes  $\frac{d}{dt}\mathbf{X}(t)$ .

One final note: rather than represent the orientation of the body as a matrix R(t) in  $\mathbf{X}(t)$ , it is better to use *quaternions*. Section 4 discusses using quaternions in place of rotation matrices. Briefly, a quaternion is a type of four element vector that can be used to represent a rotation. If we replace R(t) in  $\mathbf{X}(t)$  with a quaternion q(t), we can treat R(t) as an auxiliary variable that is computed directly from q(t), just as  $\omega(t)$  is computed from L(t). Section 4 derives a formula analogous to  $\dot{R}(t) = \omega(t)^* R(t)$ , that expresses  $\dot{q}(t)$  in terms of q(t) and  $\omega(t)$ .

# **3** Computing $\frac{d}{dt}\mathbf{X}(t)$

Lets consider an implementation of the function Dxdt() for rigid bodies. The code is written in C++, and we'll assume that we have datatypes (classes) called matrix and triple which implement, respectively,  $3 \times 3$  matrices and points in 3-space. Using these datatypes, we'll represent

```
struct RigidBody {
    /* Constant quantities */
    double mass; /* mass M */
           Ibody, /* I_{body} */
Ibodyinv; /* I_{body}^{-1} (inverse of I_{body}) */
    matrix Ibody,
    /* State variables */
                           /* x(t) */
    triple x;
                          /* R(t) */
/* P(t) */
    matrix R;
    triple P,
                            /* L(t) */
            L;
    /* Derived quantities (auxiliary variables) */
    matrix linv; /* I^{-1}(t) */
                           /* v(t) */
    triple v,
            omega; /* \omega(t) */
    /* Computed quantities */
    triple force, /* F(t) */ torque; /* \tau(t) */
};
```

and assume a global array of bodies

```
RigidBody Bodies[NBODIES];
```

The constant quantities mass, Ibody and Ibodyinv are assumed to have been calculated for each member of the array Bodies, before simulation begins. Also, the initial conditions for each rigid body are specified by assigning values to the state variables x, R, P and L of each member of Bodies. The implementation in this section represents orientation with a rotation matrix; section 4 describes the changes necessary to represent orientation by a quaternion.

We communicate with the differential equation solver ode by passing arrays of real numbers. Several bookkeeping routines are required:

```
*y++ = rb->P[0];
        *y++ = rb->P[1];
        *y++ = rb->P[2];
        *y++ = rb->L[0];
        *y++ = rb->L[1];
        *y++ = rb->L[2];
   }
and
   /* Copy information from an array into the state variables */
   void ArrayToState(RigidBody *rb, double *y)
        rb->x[0] = *y++;
        rb->x[1] = *y++;
        rb->x[2] = *y++;
        for(int i = 0; i < 3; i++)
             for(int j = 0; j < 3; j++)
                  rb->R[i,j] = *y++;
        rb - > P[0] = *y + +;
        rb - > P[1] = *y + +;
        rb - > P[2] = *y + +;
        rb->L[0] = *y++;
        rb->L[1] = *y++;
        rb->L[2] = *y++;
        /* Compute auxiliary variables... */
        \label{eq:vt} \begin{array}{ll} / * & v(t) = \frac{P(t)}{M} * / \\ \text{rb->v = rb->P / mass;} \end{array}
        /* I^{-1}(t) = R(t)I_{body}^{-1}R(t)^{T}*/
        rb->Iinv = R * Ibodyinv * Transpose(R);
        /* \omega(t) = I^{-1}(t)L(t) */
        rb->omega = rb->linv * rb->L;
   }
```

Note that ArrayToState is responsible for computing values for the auxiliary variables Iinv, v and omega. We'll assume that the appropriate arithmetic operations have been defined between real numbers, triple's and matrix's, and that Transpose returns the transpose of a matrix.

Examining these routines, we see that each rigid body's state is represented by 3+9+3+3=18 numbers. Transfers between all the members of Bodies and an array y of size  $18 \cdot \text{NBODIES}$  are implemented as

computes the force F(t) and torque  $\tau(t)$  acting on the rigid body \*rb at time t, and stores F(t) and  $\tau(t)$  in rb->force and rb->torque respectively. ComputeForceAndTorque takes into account all forces and torques: gravity, wind, interaction with other bodies etc. Using this routine, we'll define Dxdt() as

The numerical solver ode calls calls Dxdt() and is responsible for allocating enough space for the arrays y, and xdot (STATE\_SIZE·NBODIES worth for each). The function which does the real work of computing  $\frac{d}{dt}\mathbf{X}(t)$  and storing it in the array xdot is ddtStateToArray:

```
void DdtStateToArray(RigidBody *rb, double *xdot)
     / * \operatorname{copy} \ \tfrac{d}{dt} x(t) = v(t) \ \operatorname{into} \ \operatorname{xdot} \ * /
     *xdot++ = rb->v[0];
     *xdot++ = rb->v[1];
     *xdot++ = rb->v[2];
     /* Compute \dot{R}(t) = \omega(t)^* R(t) */
    matrix Rdot = Star(rb->omega) * rb->R;
     /* copy \dot{R}(t) into array */
     for(int i = 0; i < 3; i++)
          for(int j = 0; j < 3; j++)
               *xdot++ = Rdot[i,j];
                                       /* \frac{d}{dt}P(t) = F(t) */
     *xdot++ = rb->force[0];
     *xdot++ = rb->force[1];
     *xdot++ = rb->force[2];
                                      /* \frac{d}{dt}L(t) = \tau(t) */
     *xdot++ = rb->torque[0];
     *xdot++ = rb->torque[1];
     *xdot++ = rb->torque[2];
}
```

The routine Star, used to calculate  $\dot{R}(t)$  is defined as

```
matrix Star(triple a);
```

and returns the matrix

$$\begin{pmatrix} 0 & -a[2] & a[1] \\ a[2] & 0 & -a[0] \\ -a[1] & a[0] & 0 \end{pmatrix}.$$

Given all of the above, actually performing a simulation is simple. Assume that the state variables of all NBODIES rigid bodies are initialized by a routine InitStates. We'll have our simulation run for 10 seconds, calling a routine DisplayBodies every  $\frac{1}{24}$ th of a second to display the bodies:

```
for(double t = 0; t < 10.0; t += 1./24.)
{
    /* copy xFinal back to x0 */
    for(int i = 0; i < STATE_SIZE * NBODIES; i++)
    {
        x0[i] = xFinal[i];

    ode(x0, xFinal, STATE_SIZE * NBODIES,
        t, t+1./24., Dxdt);

    /* copy d/dtX(t + 1/24) into state variables */
    ArrayToBodies(xFinal);
    DisplayBodies();
}
</pre>
```

# 4 Quaternions vs. Rotation Matrices

There is a better way to represent the orientation of a rigid body than using a  $3 \times 3$  rotation matrix. For a number of reasons, *unit quaternions*, a type of four element vector normalized to unit length, are a better choice than rotation matrices[16].

For rigid body simulation, the most important reason to avoid using rotation matrices is because of numerical drift. Suppose that we keep track of the orientation of a rigid body according to the formula

$$\dot{R}(t) = \omega(t)^* R(t).$$

As we update R(t) using this formula (that is, as we integrate this equation), we will inevitably encounter drift. Numerical error will build up in the coefficients of R(t) so that R(t) will no longer be precisely a rotation matrix. Graphically, the effect would be that applying R(t) to a body would cause a skewing effect.

This problem can be alleviated by representing rotations with unit quaternions. Since quaternions have only four parameters, there is only one extra variable being used to describe the three freedoms of the rotation. In contrast, a rotation matrix uses nine parameters to describe three degrees of freedom; therefore, the degree of redundancy is noticeably lower for quaternions than rotation matrices. As a result, quaternions experience far less drift than rotation matrices. If it does become necessary to account for drift in a quaternion, it is because the quaternion has lost its unit magnitude. This is easily correctable by renormalizing the quaternion to unit length. Because of these two properties, it is desirable to represent the orientation of a body directly as a unit quaternion q(t). We will still express angular velocity as a vector  $\omega(t)$ . The orientation matrix R(t), which is needed to compute  $I^{-1}(t)$ , will be computed as an auxiliary variable from q(t).

<sup>&</sup>lt;sup>3</sup>Any quaternion of unit length corresponds to a rotation, so quaternions deviate from representing rotations only if they lose their unit length. These notes will deal with that problem in a very simplistic way.

We will write a quaternion  $s + v_x \mathbf{i} + v_y \mathbf{j} + v_z \mathbf{k}$  as the pair

$$[s, v]$$
.

Using this notation, quaternion multiplication is

$$[s_1, v_1][s_2, v_2] = [s_1 s_2 - v_1 \cdot v_2, s_1 v_2 + s_2 v_1 + v_1 \times v_2]. \tag{4-1}$$

A rotation of  $\theta$  radians about a unit axis u is represented by the unit quaternion

$$[\cos(\theta/2), \sin(\theta/2)u].$$

In using quaternions to represent rotations, if  $q_1$  and  $q_2$  indicate rotations, then  $q_2q_1$  represents the composite rotation of  $q_1$  followed by  $q_2$ .<sup>4</sup> In a moment, we'll show how to change the routines of section 3 to handle the quaternion representation for orientation. Before we can make these changes though, we'll need a formula for  $\dot{q}(t)$ . Appendix B derives the formula

$$\dot{q}(t) = \frac{1}{2}\omega(t)q(t). \tag{4-2}$$

where the multiplication  $\omega(t)q(t)$  is a shorthand for multiplication between the quaternions  $[0, \omega(t)]$  and q(t). Note the similarity between equation (4–2) and

$$\dot{R}(t) = \omega(t)^* R(t).$$

To actually use a quaternion representation, we'll need to redefine the type RigidBody:

```
struct RigidBody {
   /* Constant quantities */
                       /* mass M */
   double mass;
   /* State variables */
   quaternion q; /* x(t) */ triple P,
                       /* P(t) */
                       /* L(t) */
         L;
   /* Derived quantities (auxiliary variables) */
   matrix linv, /* I^{-1}(t) */
         R;
                      /* R(t) */
                     /* v(t) */
   triple v,
         omega;
                      /* \omega(t) */
   /* Computed quantities */
```

<sup>&</sup>lt;sup>4</sup>This is according to the convention that the rotation of a point p by a quaternion q is  $qpq^{-1}$ . Be warned! This is *opposite* the convention for rotation in the original paper Shoemake[16], but it is in correspondence with some more recent versions of Shoemake's article. Writing a composite rotation as  $q_2q_1$  parallels our matrix notation for composition of rotations.

```
triple force, /* F(t) */ torque; /* \tau(t) */ };
```

Next, in the routine StateToArray, we'll replace the double loop

A similar change is made in ArrayToState. Also, since ArrayToState is responsible for computing the auxiliary variable  $I^{-1}(t)$ , which depends on R(t), ArrayToState must also compute R(t) as an auxiliary variable: in the section

```
/* Compute auxiliary variables... */

/* v(t) = \frac{P(t)}{M} */
rb->v = rb->P / mass;

/* I^{-1}(t) = R(t)I_{body}^{-1}R(t)^T*/
rb->\mathrm{Linv} = R * \mathrm{Ibodyinv} * \mathrm{Transpose}(R);

/* \omega(t) = I^{-1}(t)L(t) */
rb->\mathrm{mega} = rb->\mathrm{Linv} * rb->\mathrm{L};

we add the line
```

prior to computing rb->Iinv. The routine normalize returns q divided by its length; this unit length quaternion returned by normalize is then passed to QuaterionToMatrix which returns a  $3 \times 3$  rotation matrix. Given a quaternion q = [s, v], QuaterionToMatrix returns the

matrix

$$\begin{pmatrix} 1 - 2v_y^2 - 2v_z^2 & 2v_x v_y - 2s v_z & 2v_x v_z + 2s v_y \\ 2v_x v_y + 2s v_z & 1 - 2v_x^2 - 2v_z^2 & 2v_y v_z - 2s v_x \\ 2v_x v_z - 2s v_y & 2v_y v_z + 2s v_x & 1 - 2v_x^2 - 2v_y^2 \end{pmatrix}.$$

In case you need to convert from a rotation matrix to a quaternion,

```
quaternion matrixToQuaternion(const matrix &m)
    quaternion
               q;
    double
             tr, s;
    tr = m[0,0] + m[1,1] + m[2,2];
    if(tr >= 0)
        s = sqrt(tr + 1);
        q.r = 0.5 * s;
        s = 0.5 / s;
        q.i = (m[2,1] - m[1,2]) * s;
        q.j = (m[0,2] - m[2,0]) * s;
        q.k = (m[1,0] - m[0,1]) * s;
    }
    else
    {
        int i = 0;
        if(m[1,1] > m[0,0])
            i = 1;
        if(m[2,2] > m[i,i))
            i = 2;
        switch (i)
        case 0:
            s = sqrt((m[0,0] - (m[1,1] + m[2,2])) + 1);
            q.i = 0.5 * s;
            s = 0.5 / s;
            q.j = (m[0,1] + m[1,0]) * s;
            q.k = (m[2,0] + m[0,2]) * s;
            q.r = (m[2,1] - m[1,2]) * s;
            break;
        case 1:
            s = sqrt((m[1,1] - (m[2,2] + m[0,0])) + 1);
            q.j = 0.5 * s;
            s = 0.5 / s;
```

```
q.k = (m[1,2] + m[2,1]) * s;
q.i = (m[0,1] + m[1,0]) * s;
q.r = (m[0,2] - m[2,0]) * s;
break;
case 2:
    s = sqrt((m[2,2] - (m[0,0] + m[1,1])) + 1);
q.k = 0.5 * s;
s = 0.5 / s;
q.i = (m[2,0] + m[0,2]) * s;
q.j = (m[1,2] + m[2,1]) * s;
q.r = (m[1,0] - m[0,1]) * s;
}
return q;
}
```

The matrix m is structured so that m[0,0], m[0,1] and m[0,2] form the first row (not column) of m.

The routines ArrayToBodies and BodiesToArray don't need any changes at all, but note that the constant STATE\_SIZE changes from 18 to 13, since a quaternion requires five less elements than a rotation matrix. The only other change we need is in ddtStateToArray. Instead of

We're assuming here that the multiplication between the triple rb->omega and the quaternion rb->q is defined to return the quaternion product

```
[0, rb->omega]q.
```

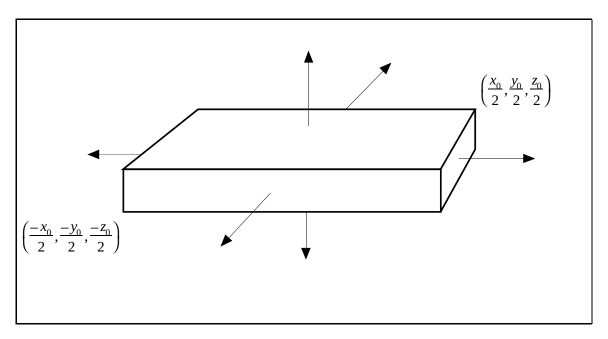


Figure 7: A rectangular block of constant unit density, with center of mass at (0,0,0).

# 5 Examples

#### 5.1 Inertia Tensor of a Block

Let us calculate the inertia tensor  $I_{body}$  of the rectangular block in figure 7. The block has dimensions  $x_0 \times y_0 \times z_0$ . As required, the center of mass of the block is at the origin. Thus, the extent of the block is from  $-\frac{x_0}{2}$  to  $\frac{x_0}{2}$  along the x axis, and similarly for the y and z axes. To calculate the inertia tensor, we must treat the sums in equation (2–32) as integrals over the volume of the block. Let us assume that the block has constant unit density. This means that the density function  $\rho(x, y, z)$  is always one. Since the block has volume  $x_0y_0z_0$ , the mass M of the block is  $M = x_0y_0z_0$ . Then, in

body space,

$$I_{xx} = \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \int_{-\frac{y_0}{2}}^{\frac{y_0}{2}} \int_{-\frac{z_0}{2}}^{\frac{z_0}{2}} \rho(x, y, z) (y^2 + z^2) dx dy dz = \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \int_{-\frac{y_0}{2}}^{\frac{y_0}{2}} \int_{-\frac{z_0}{2}}^{\frac{z_0}{2}} y^2 + z^2 dx dy dz$$

$$= \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \int_{-\frac{y_0}{2}}^{\frac{y_0}{2}} y^2 z + \frac{z^3}{3} \Big|_{z=-\frac{z_0}{2}}^{z=-\frac{z_0}{2}} dx dy$$

$$= \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \int_{-\frac{y_0}{2}}^{\frac{y_0}{2}} y^2 z_0 + \frac{z_0^3}{12} dx dy$$

$$= \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \frac{y_0^3}{3} z_0 + \frac{z_0^3}{12} y_0 \Big|_{y=-\frac{y_0}{2}}^{y=\frac{y_0}{2}} dx$$

$$= \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \frac{y_0^3 z_0}{12} + \frac{z_0^3 y_0}{12} dx$$

$$= \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \frac{y_0^3 z_0}{12} + \frac{z_0^3 y_0}{12} dx$$

$$= \frac{y_0^3 z_0}{12} + \frac{z_0^3 y_0}{12} \Big|_{x=-\frac{x_0}{2}}^{x=-\frac{x_0}{2}} \frac{y_0^3 z_0 x_0}{12} + \frac{z_0^3 y_0 x_0}{12} = \frac{x_0 y_0 z_0}{12} (y_0^2 + z_0^2) = \frac{M}{12} (y_0^2 + z_0^2).$$

Similarly,  $I_{yy} = \frac{M}{12}(x_0^2 + z_0^2)$  and  $I_{zz} = \frac{M}{12}(x_0^2 + y_0^2)$ . Now, the off-diagonal terms, such as  $I_{xy}$ , are

$$I_{xy} = \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \int_{-\frac{y_0}{2}}^{\frac{y_0}{2}} \int_{-\frac{z_0}{2}}^{\frac{z_0}{2}} \rho(x, y, z)(xy) \, dx \, dy \, dz = \int_{-\frac{x_0}{2}}^{\frac{x_0}{2}} \int_{-\frac{y_0}{2}}^{\frac{y_0}{2}} \int_{-\frac{z_0}{2}}^{\frac{z_0}{2}} xy \, dx \, dy \, dz = 0$$
 (5-2)

(and similarly for the others) because the integrals are all symmetric. Thus, the inertia tensor of the block is

$$I_{body} = \frac{M}{12} \begin{pmatrix} y_0^2 + z_0^2 & 0 & 0\\ 0 & x_0^2 + z_0^2 & 0\\ 0 & 0 & x_0^2 + y_0^2 \end{pmatrix}.$$
 (5-3)

#### 5.2 A Uniform Force Field

Suppose a uniform force acts on each particle of a body. For example, we typically describe a gravitational field as exerting a force  $m_i g$  on each particle of a rigid body, where g is a vector pointing downwards. The net force  $F_g$  acting due to gravity on the body then is

$$F_g = \sum m_i g = Mg \tag{5-4}$$

which yields an acceleration of  $\frac{Mg}{g} = g$  of the center of mass, as expected. What is the torque due to the gravitational field? The net torque is the sum

$$\sum (r_i(t) - x(t)) \times m_i g = \left(\sum m_i (r_i(t) - x(t))\right) \times g = \mathbf{0}$$
 (5-5)

by equation (2-20). We see from this that a uniform gravitational field can have no effect on the angular momentum of a body. Furthermore, the gravitational field can be treated as a single force Mg acting on the body at its center of mass.

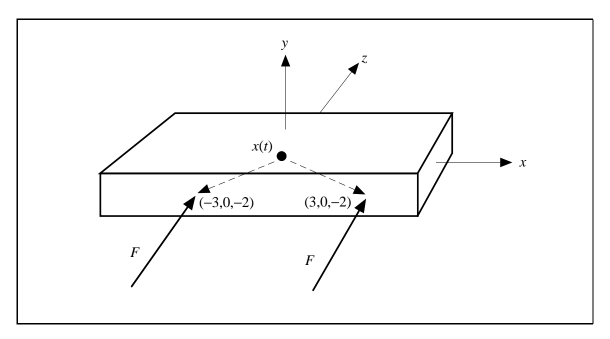


Figure 8: A block acted on by two equal forces F at two different points.

#### 5.3 Rotation Free Movement of a Body

Now, let us consider some forces acting on the block of figure 8. Suppose that an external force F = (0, 0, f) acts on the body at points x(t) + (-3, 0, -2) and x(t) + (3, 0, -2). We would expect that this would cause the body to accelerate linearly, without accelerating angularly. The net force acting on the body is (0, 0, 2f), so the acceleration of the center of mass is

$$\frac{2f}{M}$$

along the z axis. The torque due to the force acting at x(t) + (-3, 0, -2) is

$$((x(t) + \begin{pmatrix} -3 \\ 0 \\ -2 \end{pmatrix}) - x(t)) \times F = \begin{pmatrix} -3 \\ 0 \\ -2 \end{pmatrix} \times F$$

while the torque due to the force acting at x(t) + (3, 0, -2) is

$$((x(t) + \begin{pmatrix} 3 \\ 0 \\ -2 \end{pmatrix}) - x(t)) \times F = \begin{pmatrix} 3 \\ 0 \\ -2 \end{pmatrix} \times F.$$

The total torque  $\tau$  is therefore

$$\tau = \begin{pmatrix} -3 \\ 0 \\ -2 \end{pmatrix} \times F + \begin{pmatrix} 3 \\ 0 \\ -2 \end{pmatrix} \times F = \begin{pmatrix} -3 \\ 0 \\ -2 \end{pmatrix} + \begin{pmatrix} 3 \\ 0 \\ -2 \end{pmatrix}) \times F = \begin{pmatrix} 0 \\ 0 \\ -2 \end{pmatrix} \times F.$$

But this gives

$$\tau = \begin{pmatrix} 0 \\ 0 \\ -2 \end{pmatrix} \times \begin{pmatrix} 0 \\ 0 \\ f \end{pmatrix} = \mathbf{0}.$$

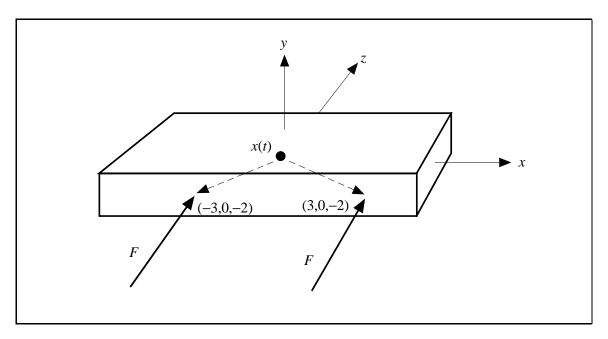


Figure 9: A block acted on by two opposite forces  $F_1$  and  $F_2 = -F_1$ , at two different points.

As expected then, the forces acting on the block impart no angular acceleration to the block.

## 5.4 Translation Free Movement of a Body

Suppose now that an external force  $F_1 = (0, 0, f)$  acts on the body at point x(t) + (-3, 0, -2) and an external force  $F_2 = (0, 0, -f)$  acts on the body at point x(t) + (3, 0, 2) (figure 9). Since  $F_1 = -F_2$ , the net force acting on the block is  $F_1 + F_2 = \mathbf{0}$ , so there is no acceleration of the center of mass. On the other hand, the net torque is

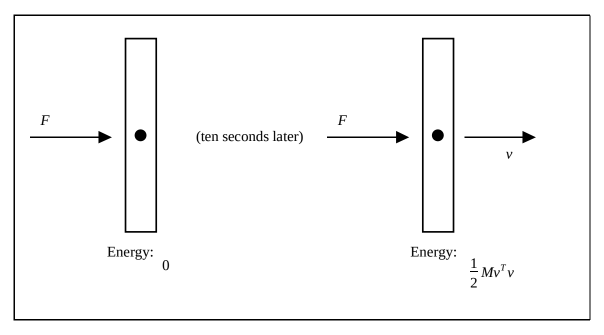


Figure 10: A rectangular block acted on by a force through its center of mass.

$$((x(t) + \begin{pmatrix} -3 \\ 0 \\ -2 \end{pmatrix}) - x(t)) \times F_1 +$$

$$((x(t) + \begin{pmatrix} 3 \\ 0 \\ 2 \end{pmatrix}) - x(t)) \times F_2 = \begin{pmatrix} -3 \\ 0 \\ -2 \end{pmatrix} \times \begin{pmatrix} 0 \\ 0 \\ f \end{pmatrix} + \begin{pmatrix} 3 \\ 0 \\ -2 \end{pmatrix} \times \begin{pmatrix} 0 \\ 0 \\ -f \end{pmatrix}$$

$$= \begin{pmatrix} 0 \\ 3f \\ 0 \end{pmatrix} + \begin{pmatrix} 0 \\ 3f \\ 0 \end{pmatrix} = \begin{pmatrix} 0 \\ 6f \\ 0 \end{pmatrix}.$$

$$(5-6)$$

Thus, the net torque is (0, 6f, 0), which is parallel to the y axis. The final result is that the forces acting on the block cause it to angularly accelerate about the y axis.

#### 5.5 Force vs. Torque Puzzle

In considering the effect of a force acting at a point on a body, it sometimes seems that the force is being considered twice. That is, if a force F acts on a body at a point r + x(t) in space, then we first consider F as accelerating the center of mass, and then consider F as imparting a spin to the body.

This gives rise to what at first seems a paradox: Consider the long horizontal block of figure 10 which is initially at rest. Suppose that a force F acts on the block at the center of mass for some period of time, say, ten seconds. Since the force acts at the center of mass, no torque is exerted on the body. After ten seconds, the body will have acquired some linear velocity v. The body will not have acquired any angular velocity; thus the kinetic energy of the block will be  $\frac{1}{2}M|v|^2$ .

Now suppose that the same force F is applied off-center to the body as shown in figure 11. Since the force acting on the body is the same, the acceleration of the center of mass is the same. Thus,

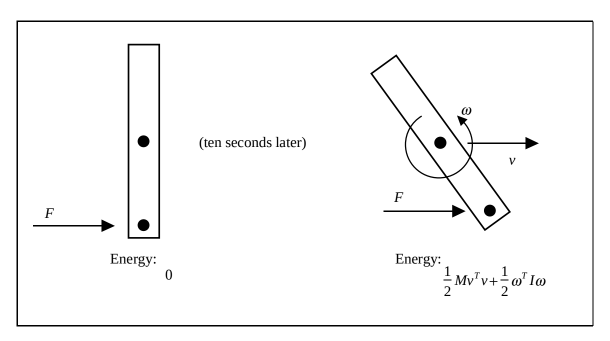


Figure 11: A block acted on by a force, off-center of the center of mass.

after ten seconds, the body will again have linear velocity v. However, after ten seconds, the body will have picked up some angular velocity  $\omega$ , since the force F, acting off center, now exerts a torque on the body. Since the kinetic energy is (see appendix C)

$$\frac{1}{2}M|v|^2 + \frac{1}{2}\omega^T I\omega$$

the kinetic energy of the block is higher than when the force acted through the center of mass. But if identical forces pushed the block in both cases, how can the energy of the block be different? Hint: Energy, or work, is the integral of force over distance.

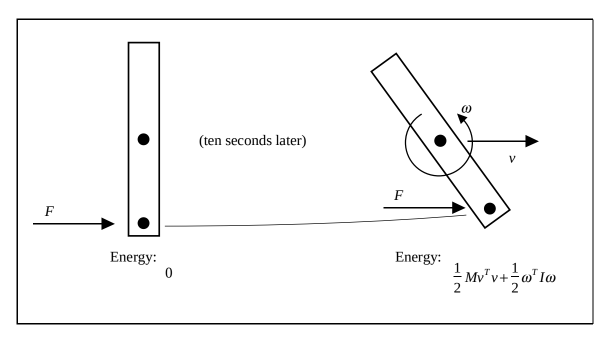


Figure 12: The path the force acts over is longer than in figure 10. As a result, the force does more work, imparting a larger kinetic energy to the block.

Figure 12 shows why the force acting off center results in a higher kinetic energy. The kinetic energy of the block is equivalent to the work done by the force. The work done by the force is the integral of the force over the path traveled in applying that force. In figure 11, where the force acts off the center of mass, consider the path traced out by the point where the force is applied. This path is clearly longer than the path taken by the center of mass in figure 10. Thus, when the force is applied off center, more work is done because the point p at which the force is applied traces out a longer path then when the force is applied at the center of mass.

# **Part II. Nonpenetration Constraints**

## **6** Problems of Nonpenetration Constraints

Now that we know how to write and implement the equations of motion for a rigid body, let's consider the problem of preventing bodies from inter-penetrating as they move about an environment. For simplicity, suppose we simulate dropping a point mass (i.e. a single particle) onto a fixed floor. There are several issues involved here.

Because we are dealing with rigid bodies, that are totally non-flexible, we don't want to allow any inter-penetration at all when the particle strikes the floor. (If we considered our floor to be flexible, we might allow the particle to inter-penetrate some small distance, and view that as the floor actually deforming near where the particle impacted. But we don't consider the floor to be flexible, so we don't want any inter-penetration at all.) This means that at the instant that the particle actually comes into contact with the floor, what we would like is to abruptly change the velocity of the particle. This is quite different from the approach taken for flexible bodies. For a flexible body, say a rubber ball, we might consider the collision as occurring gradually. That is, over some fairly small, but non-zero span of time, a force would act between the ball and the floor and change the ball's velocity. During this time span, the ball would deform, due to the force. The more rigid we made the ball, the less the ball would deform, and the faster this collision would occur. In the limiting case, the ball is infinitely rigid, and can't deform at all. Unless the ball's downward velocity is halted instantaneously, the ball will inter-penetrate the floor somewhat. In rigid body dynamics then, we consider collisions as occurring instantaneously.

This means we have two types of contact we need to deal with. When two bodies are in contact at some point p, and they have a velocity towards each other (as in the particle striking the floor), we call this *colliding contact*. Colliding contact requires an instantaneous change in velocity. Whenever a collision occurs, the state of a body, which describes both position, and velocity, undergoes a discontinuity in the velocity. The numerical routines that solve ODE's do so under the assumption that the state  $\mathbf{X}(t)$  always varies smoothly. Clearly, requiring  $\mathbf{X}(t)$  to change discontinuously when a collision occurs violates that assumption.

We get around this problem as follows. If a collision occurs at time  $t_c$ , we tell the ODE solver to stop. We then take the state at this time,  $\mathbf{X}(t_c)$ , and compute how the velocities of bodies involved in the collision must change. We'll call the state reflecting these new velocities  $\mathbf{X}(t_c)^+$ . Note that  $\mathbf{X}(t_c)$  and  $\mathbf{X}(t_c)^+$  agree for all spatial variables (position and orientation), but will be different for the velocity variables of bodies involved in the collision at time  $t_c$ . We then restart the numerical solver, with the new state  $\mathbf{X}(t_c)$ , and instruct it to simulate forward from time  $t_c$ .

Whenever bodies are resting on one another at some point p (e.g. imagine the particle in contact with the floor with zero velocity), we say that the bodies are in *resting contact*. In this case, we compute a force that prevents the particle from accelerating downwards; essentially, this force is the weight of the particle due to gravity (or whatever other external forces push on the particle). We call the force between the particle and the floor a *contact force*. Resting contact clearly doesn't require us to stop and restart the ODE solve at every instant; from the ODE solver's point of view, contact forces are just a part of the force returned by ComputeForceAndTorque.

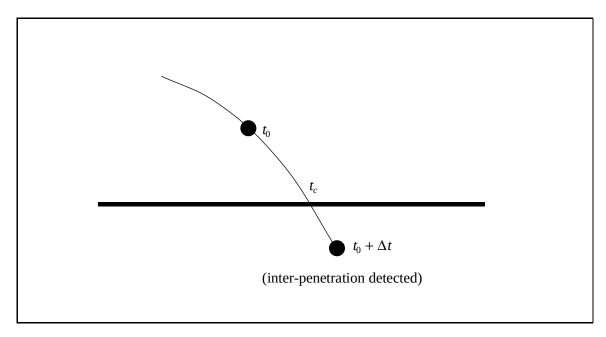


Figure 13: At time  $t_0 + \Delta t$ , the particle is found to lie below the floor. Thus, the actual time of collision  $t_c$  lies between the time of the last known legal position,  $t_0$ , and  $t_0 + \Delta t$ .

So far then, we have two problems we'll need to deal with: computing velocity changes for *colliding contact*, and computing the contact forces that prevent inter-penetration. But before we can tackle these problems we have to deal with the geometric issue of actually detecting contact between bodies. Let's go back to dropping the particle to the floor. As we run our simulation, we compute the position of the particle as it drops towards the floor at specific time values (figure 13). Suppose we consider the particle at times  $t_0$ ,  $t_0 + \Delta t$ ,  $t_0 + 2\Delta t$  etc.<sup>5</sup> and suppose the time of collision,  $t_c$ , at which the particle actually strikes the floor, lies between  $t_0$  and  $t_0 + \Delta t$ . Then at time  $t_0$ , we find that the particle lies above the floor, but at the next time step,  $t_0 + \Delta t$ , we find the particle is beneath the floor, which means that inter-penetration has occurred.

If we're going to stop and restart the simulator at time  $t_c$ , we'll need to compute  $t_c$ . All we know so far is that  $t_c$  lies between  $t_0$  and  $t_0 + \Delta t$ . In general, solving for  $t_c$  exactly is difficult, so we solve for  $t_c$  numerically, to within a certain tolerance. A simple way of determining  $t_c$  is to use a numerical method called bisection[14]. If at time  $t_0 + \Delta t$  we detect inter-penetration, we inform the ODE solver that we wish to restart back at time  $t_0$ , and simulate forward to time  $t_0 + \Delta t/2$ . If the simulator reaches  $t_0 + \Delta t/2$  without encountering inter-penetration, we know the collision time  $t_c$  lies between  $t_0 + \Delta t/2$  and  $t_0 + \Delta t$ . Otherwise,  $t_c$  is less than  $t_0 + \Delta t/2$ , and we try to simulate from  $t_0$  to  $t_0 + \Delta t/4$ . Eventually, the time of collision  $t_c$  is computed to within some suitable numerical tolerance. The accuracy with which  $t_c$  is found depends on the collision detection routines. The collision detection routines have some parameter  $\epsilon$ . We decide that our computation of  $t_c$  is "good enough" when the particle inter-penetrates the floor by no more than  $\epsilon$ , and is less than  $\epsilon$  above the floor. At this point we declare that the particle is in contact with the floor (figure 14).

The method of bisection is a little slow, but its easy to implement and quite robust. A faster method involves actually predicting the time  $t_c$  of the collision, based on examining  $\mathbf{X}(t_0)$  and  $\mathbf{X}(t_0 + \Delta t)$ . Baraff[1, 2] describes how to make such predictions. How to actually implement all of

<sup>&</sup>lt;sup>5</sup>The ODE solver doesn't have to proceed with equal size time steps though.

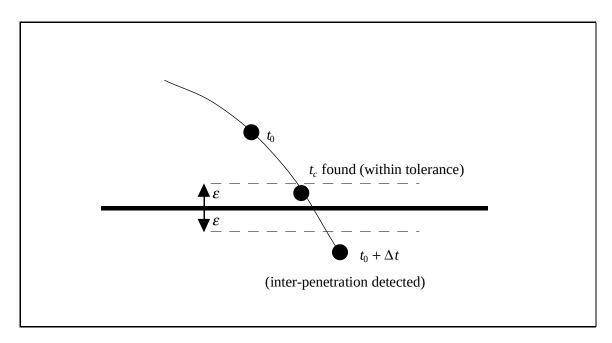


Figure 14: When the particle is found to be within some tolerance  $\epsilon$  of contacting the floor, then  $t_c$  is considered to have been computed to within sufficient accuracy.

this depends on how you interact with your ODE routines. One might use exception handling code to signal the ODE of various events (collisions, inter-penetration), or pass some sort of messages to the ODE solver. We'll just assume that you have some way of getting your ODE solver to progress just up to the point  $t_c$ .

Once you actually reach the time of a collision, or whenever you're in a state  $\mathbf{X}(t)$  where no inter-penetration has occurred, a geometric determination has to be made to find all the points of contact. (Just because you may be looking for the time of collision between two bodies A and B doesn't mean you get to neglect resting contact forces between other bodies C and D. Whenever you're trying to move the simulation forward, you'll need to compute the point of contact between bodies and the contact forces at those points.) There is a vast amount of literature dealing with the collision detection problem. For instance, some recent SIGGRAPH papers dealing with the subject are Von Herzen, Barr and Zatz[17] and Moore and Wilhelms[12]; in robotics, a number of papers of interest are Canny[4], Gilbert and Hong[6], Meyer[11] and Cundall[5]. Preparata and Shamos[13] describes many approaches in computational geometry to the problem. In the next section, we'll briefly describe a collision detection "philosophy" that leads to very efficient algorithms, for the sorts of simulation these course notes are concerned with. Actual code for the algorithms is fairly easy to write, but a little too lengthy to fit in these notes. Following this, we'll move on to consider colliding and resting contact.

# 7 Collision Detection

The collision detection algorithm begins with a preprocessing step, in which a bounding box for each rigid body is computed (a box with sides parallel to the coordinate axes). Given n such bounding boxes, we will want to quickly determine all pairs of bounding boxes that overlap. Any pair of rigid bodies whose bounding boxes do not overlap need not be considered any further. Pairs of rigid

bodies whose bounding boxes do overlap require further consideration. We'll first describe how to efficiently check for inter-penetration or contact points between rigid bodies defined as convex polyhedra. Then we'll show how to perform the bounding box check efficiently.

As described in section 1, the simulation process consists of the repeated computation of the derivative of the state vector,  $\frac{d}{dt}\mathbf{X}(t)$ , at various times t. The numerical ODE solver is responsible for choosing the values of t at which the state derivative is to be computed. For any reasonably complicated simulation, the values of t chosen are such that the state  $\mathbf{X}$  does not change greatly between successive values of t. As a result, there is almost always great geometric coherence between successive time steps. At a time step  $t_0 + \Delta t$ , the idea is to take advantage of the collision detection results computed at the previous time step  $t_0$ .

#### 7.1 Convex Polyhedra

Our primary mechanism for exploiting coherence will be through the use of *witnesses*. In our context, given two convex polyhedra A and B, a witness is some piece of information that can be used to quickly answer the "yes/no" question "are A and B disjoint"? We will utilize coherence by caching witnesses from one time step to the next; hopefully a witness from the previous time step will be a witness during the current time step.

Since we are considering convex polyhedra, two polyhedra do not inter-penetrate if and only if a separating plane between them exists. A separating plane between two polyhedra is a plane such that each polyhedron lies on a different side of the plane. A given plane can be verified to be a separating plane by testing to make sure that all of the vertices of *A* and *B* lie on opposite sides of the plane. Thus, a separating plane is a witness to the fact that two convex polyhedra do not inter-penetrate. If a separating plane does not exist, then the polyhedra must be inter-penetrating.

The cost of initially finding a witness (for the very first time step of the simulation, or the first time two bodies become close enough to require more than a bounding box test) is unavoidable. A simple way to find a separating plane initially is as follows. If a pair of convex polyhedra are disjoint or contacting (but not inter-penetrating), then a separating plane exists with the following property: either the plane contains a face of one of the polyhedra, or the plane contains an edge from one of the polyhedra and is parallel to an edge of the other polyhedra. (That is, the separating plane's normal is the cross product of the two edge directions, and the plane itself contains one of the edges.) We will call the face or edges in question the *defining* face or edges. Initially, we simply check all possible combinations of faces and edges to see if one such combination forms a separating plane (figure 15). Although this is inefficient, it's done so infrequently that the inefficiency is unimportant. For subsequent time steps, all we need to do is form a separating plane from the defining face or edges found during the previous time step, and then verify the plane to see that it is still valid (figure 16).

On those (rare) occasions when the cached face or two edges fails to form a valid separating plane (figure 17), faces or edges adjacent to the previously cached face or edges can be examined to see if they form a separating plane; however, this happens infrequently enough that it may be simpler to start from scratch and compute a new separating plane without using any prior knowledge.

Once the separating place has been found, the contact region between the two polyhedra is determined, assuming the polyhedra are not disjoint. Contact points between the two polyhedra can only occur on the separating plane. Given the separating plane, the contact points can be quickly and efficiently determined by comparing only those faces, edges, and vertices of the polyhedra that are coincident with the separating plane.

However, if no separating plane can be found, then the two polyhedra must be inter-penetrating. When two polyhedra inter-penetrate, it is almost always the case that either a vertex of one poly-

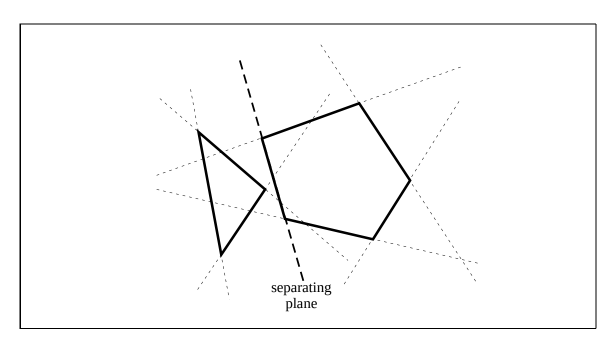


Figure 15: Exhaustive search for a separating plane. Only one face of the two polygons forms a separating plane.

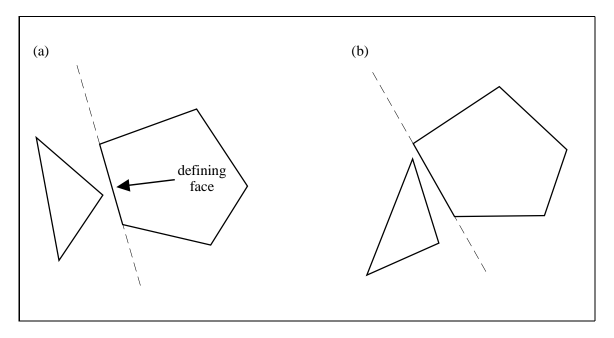


Figure 16: (a) At this time step, the separating plane is defined by a face of one of the polygons. (b) At the next time step, the polygons have moved, but the same face still defines a separating plane.

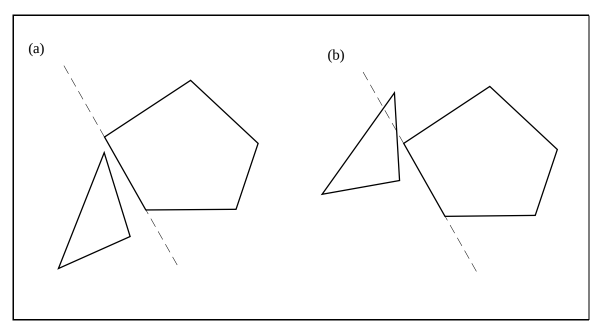


Figure 17: The face that has been defining a separating plane no longer does so, and a new separating plane must be found.

hedron is inside the other, or an edge of one polyhedron has intersected a face of the other.<sup>6</sup> In this case, the inter-penetrating vertex, or intersecting edge and face are cached as a witness to the inter-penetration. Since this indicates a collision at some earlier time, the simulator will back up and attempt to compute  $\frac{d}{dt}\mathbf{X}(t)$  at some earlier time. Until the collision time is determined, the first action taken by the collision/contact determination step will be to check the cached vertex or edge and face to see if they indicate inter-penetration. Thus, until the collision time is found, states in which the inter-penetration still exists are identified as such with a minimum of computational overhead.

#### 7.2 Bounding Boxes

To reduce the number of pairwise collision/contact determinations necessary, a bounding box hierarchy is imposed on the bodies in the simulation environment. If two bounding boxes are found not to overlap, no further comparisons involving the contents of the boxes are needed. Given a collection of n rectangular bounding boxes, aligned with the coordinate axes, we would like to efficiently determine all pairs of boxes that overlap. A naive pairwise comparison of all pairs requires  $O(n^2)$  work and is too inefficient, unless the number of bodies is small. Computational geometry algorithms exist that can solve this problem in time O(nlogn + k) where k is the number of pairwise overlaps; a general result is that the problem can be solved in time  $O(nlog^{d-2}n + k)$  for d-dimensional bounding boxes[13]. Using coherence, we can achieve substantially better performance.

<sup>&</sup>lt;sup>6</sup>An exception is the following. Stack two cubes of equal size atop one another so that their contacting faces exactly coincide. Lower the top one. This produces an inter-penetration such that no vertex is inside either cube, and no edge penetrates through any face.

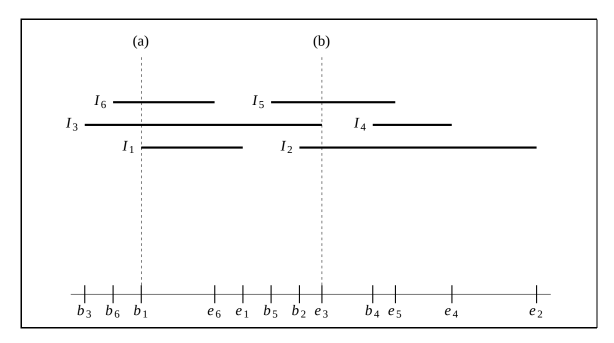


Figure 18: The sweep/sort algorithm. (a) When  $b_1$  is encountered, the active list contains intervals 3 and 6; interval 1 is reported to overlap with these two intervals. Interval 1 is added to the active list and the algorithm continues. (b) When  $e_3$  is encountered, the active list contains intervals 2, 3 and 5. Interval 3 is removed from the active list.

#### 7.2.1 The one-dimensional case

Consider the problem of detecting overlap between *one*-dimensional bounding boxes, aligned with the coordinate system. Such a bounding box can be described simply as an interval [b, e] where b and e are real numbers. Let us consider a list of n such intervals, with the ith interval being  $[b_i, e_i]$ . The problem is then defined to be the determination of all pairs i and j such that the intervals  $[b_i, e_i]$  and  $[b_i, e_j]$  intersect.

The problem can be solved initially by a *sort and sweep* algorithm. A sorted list of all the  $b_i$  and  $e_i$  values is created, from lowest to highest. The list is then swept, and a list of *active* intervals, initially empty, is maintained. Whenever some value  $b_i$  is encountered, all intervals on the active list are output as overlapping with interval i, and interval i is then added to the list (figure 18a). Whenever some value  $e_i$  is encountered, interval i is removed from the active list (figure 18b). The cost of this process is  $O(n \log n)$  to create the sorted list, O(n) to sweep through the list, and O(k) to output each overlap. This gives a total cost of  $O(n \log n + k)$ , and is an optimal algorithm for initially solving the problem.

Subsequent comparisons can be improved as follows. First, there is no need to use an  $O(n \log n)$  algorithm to form the sorted list of  $b_i$  and  $e_i$  values. It is considerably more efficient to start with the order found for  $b_i$  and  $e_i$  values from the previous time step; if coherence is high, this ordering will be nearly correct for the current time step. A sorting method called an *insertion sort*[15] is used to permute the "nearly sorted" list into a sorted list. The insertion sort algorithm works by moving items towards the beginning of the list, until a smaller item is encountered. Thus, the second item is interchanged with the first if necessary, then the third item is moved towards the beginning of the list until its proper place is found, and so on; each movement of an item indicates a change in the ordering of two values. After the last item on the list has been processed, the list is in order. Such

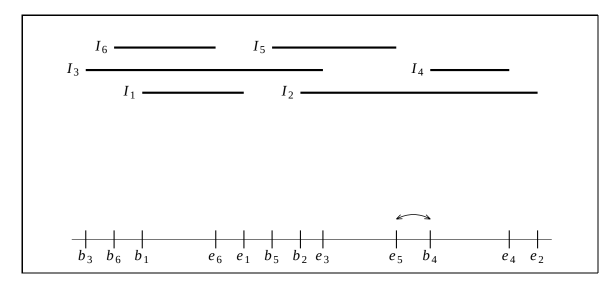


Figure 19: A coherence-based method of detecting overlaps. The order produced in figure 18 is nearly correct for this arrangement of intervals. Only  $b_4$  and  $e_5$  need to be exchanged. When the exchange occurs, the change in overlap status between interval 4 and 5 is detected.

a sort takes time O(n+c) where c is the number of exchanges necessary. For example, the only difference between figures 19 and 18 is that interval 4 has moved to the right. Starting from the ordered list of  $b_i$  and  $e_i$  values of figure 18, only a single exchange is necessary to sort the list for figure 19. The insertion sort is not recommendeded as a sorting procedure in general, since it may require  $O(n^2)$  exchanges; however, it is a good algorithm for sorting a nearly sorted list, which is what occurs in our highly coherent environment. To complete the algorithm, note that if two intervals i and j overlap at the previous time step, but not at the current time step, one or more exchanges involving either a  $b_i$  or  $e_i$  value and a  $b_j$  or  $e_j$  value must occur. The converse is true as well when intervals i and j change from not overlapping at the previous time step to overlapping at the current time step.

Thus, if we maintain a table of overlapping intervals at each time step, the table can be updated at each time step with a total cost of O(n+c). Assuming coherence, the number of exchanges c necessary will be close to the actual number k of changes in overlap status, and the extra O(c-k) work will be negligible. Thus, for the one-dimensional bounding box problem, the coherence view yields an efficient algorithm of extreme (if not maximal) simplicity that approaches optimality as coherence increases.

#### 7.2.2 The three-dimensional case

Efficient computational geometry algorithms for solving the bounding box intersection problem in  $\mathbb{R}^3$  are much more complicated than the sort and sweep method for the one-dimensional case. However, these algorithms all have in common a step that is essentially a sort along a coordinate axis, as in the one-dimensional case. Each bounding box is described as three independent intervals  $[b_i^{(x)}, e_i^{(x)}]$ ,  $[b_i^{(y)}, e_i^{(y)}]$ , and  $[b_i^{(z)}, e_i^{(z)}]$  which represent the intervals spanned on the three coordinate axes by the *i*th bounding box. Thus, our first thought towards improving the efficiency of a computational geometry algorithm for coherent situations would be to sort a list containing the  $b_i^{(x)}$  and  $e_i^{(x)}$  values, and similarly for the *y* and *z* axes. Again, such a step will involve O(n+c) work, where *c* is now the

total number of exchanges involved in sorting all three lists. However, if we observe that checking two bounding boxes for overlap is a constant time operation, it follows that if we simply check bounding boxes i and j for overlap whenever an exchange is made between values indexed by i and j (on any coordinate axis), we will detect all changes in overlap status in O(n+c) time.

Again, we can maintain a table of overlapping bounding boxes, and update it at each time step in O(n+c) time. The extra work involved is again O(c-k). For the three-dimensional case, extra work can occur if the extents of two bounding boxes change on one coordinate axis without an actual change of their overlap status. In practice, the extra work done has been found to be completely negligible, and the algorithm runs essentially in time O(n+k).

### **8 Colliding Contact**

For the remainder of these notes, we're going to be concerned with examining the bodies in our simulator at a particular instant of time  $t_0$ . At this time  $t_0$ , we assume that no bodies are interpenetrating, and that the simulator has already determined which bodies contact, and at which points. To simplify matters, we'll imagine that all bodies are polyhedra, and that every contact point between bodies has been detected. We'll consider contacts between polyhedra as either vertex/face contacts or edge/edge contacts. A vertex/face contact occurs when a vertex on one polyhedra is in contact with a face on another polyhedra. An edge/edge contact occurs when a pair of edges contact; it is assumed in this case that the two edges are not collinear. (Vertex/vertex and vertex/edge contacts are degenerate, and are not considered in these notes.) As examples, a cube resting on a plane would be described as four vertex/face contacts, one contact at each corner of the cube. A cube resting on a table, but with its bottom face hanging over the edge of the table would still be described as four contacts; two vertex/face contacts for the vertices on the table, and two edge/edge contacts, one on each edge of the cube that crosses over an edge of the table.

Each contact is represented by a structure

```
struct Contact {
    RigidBody
                     *a,
                             /* body containing vertex */
                             /* body containing face */
                     *b;
                             /* world-space vertex location */
    triple
                     p,
                             /* outwards pointing normal of face */
                     n,
                             /* edge direction for A */
                     ea,
                             /* edge direction for B */
                     eb;
                             /* true if vertex/face contact */
    bool
                     vf;
};
int
        Ncontacts;
Contact *Contacts;
```

If the contact is a vertex/face contact, then the variable a points to the rigid body that the contact vertex is attached to, while b points to the body the face is attached to. We'll call these two bodies *A* and *B* respectively. For vertex/face contacts, the variable n is set to the outwards pointing unit normal of the contact face of body *B*, and the variables ea and eb are unused.

For edge/edge contacts, ea is a triple of unit length, that points in the direction of the contacting edge of body A (pointed to by a). Similarly, eb is a unit vector giving the direction that the contact

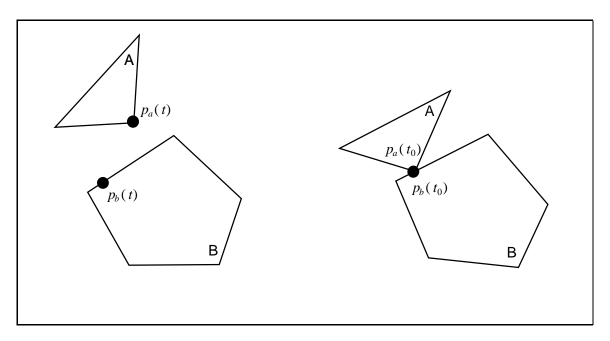


Figure 20: (a) The points  $p_a(t)$  and  $p_b(t)$  for a vertex/face contact. (b) At time  $t_0$ , the bodies come into contact at  $p_a(t_0) = p_b(t_0)$ .

edge on body B points. For edge/edge contacts, n denotes a unit vector in the ea  $\times$  eb direction. We'll adopt the convention that the two contacting bodies are labeled A and B such that the normal direction ea  $\times$  eb points outwards from B, towards A, as it does for vertex/face contacts.

For both types of contact, the position of the contact in world space (which is either the contact vertex, or the point where the two edges intersect) is given by p. The collision detection routines are responsible for discovering all the contact points, setting Ncontacts to the number of contact points, and allocating space for and initializing an array of Contact structures.

The first thing we'll need to do is examine the data in each Contact structure to see if colliding contact is taking place. For a given contact point, the two bodies A and B contact at the point p. Let  $p_a(t)$  denote the particular the point on body A that satisfies  $p_a(t_0) = p$ . (For vertex/face contacts, this point will be the vertex itself. For edge/edge contacts, it is some particular point on the contact edge of A.) Similarly, let  $p_b(t)$  denote the particular point on body B that coincides with  $p_a(t_0) = p$  at time  $t_0$  (figure 20). Although  $p_a(t)$  and  $p_b(t)$  are coincident at time  $t_0$ , the *velocity* of the two points at time  $t_0$  may be quite different. We will examine this velocity to see if the bodies are colliding or not.

From section 2.5, we can calculate the velocity of the vertex point,  $\dot{p}_a(t_0)$  by the formula

$$\dot{p}_a(t_0) = v_a(t_0) + \omega_a(t_0) \times (p_a(t_0) - x_a(t_0)) \tag{8-1}$$

where  $v_a(t)$  and  $\omega_a(t)$  are the velocities for body A. Similarly, the velocity of the contact point on the face of B is

$$\dot{p}_b(t_0) = v_b(t_0) + \omega_b(t_0) \times (p_b(t_0) - x_b(t_0)). \tag{8-2}$$

Let's examine the quantity

$$v_{rel} = \hat{n}(t_0) \cdot (\dot{p}_a(t_0) - \dot{p}_b(t_0)),$$
 (8-3)

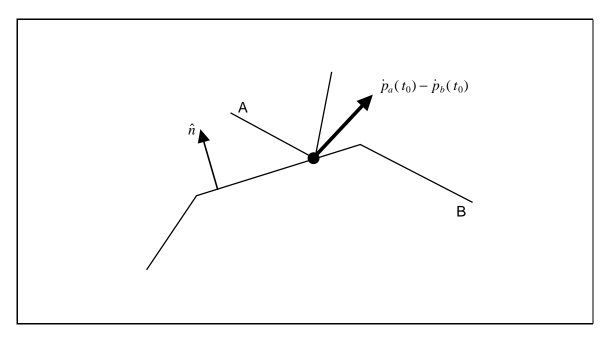


Figure 21: The vector  $\dot{p}_a(t_0) - \dot{p}_b(t_0)$  points in the same direction as  $\hat{n}(t_0)$ ; the bodies are separating.

which is a scalar. In this equation,  $\hat{n}(t_0)$  is the unit surface normal, described by the variable n, for each contact point. The quantity  $v_{rel}$  gives the component of the relative velocity  $\dot{p}_a(t_0) - \dot{p}_b(t_0)$  in the  $\hat{n}(t_0)$  direction. Clearly, if  $v_{rel}$  is positive, then the relative velocity  $\dot{p}_a(t_0) - \dot{p}_b(t_0)$  at the contact point is in the positive  $\hat{n}(t_0)$  direction. This means that the bodies are moving apart, and that this contact point will disappear immediately after time  $t_0$  (figure 21). We don't need to worry about this case. If  $v_{rel}$  is zero, then the bodies are neither approaching nor receding at p (figure 22). This is exactly what we mean by resting contact, and we'll deal with it in the next section.

In this section, we're interested in the last possibility, which is  $v_{rel} < 0$ . This means that the relative velocity at p is opposite  $\hat{n}(t_0)$ , and we have colliding contact. If the velocities of the bodies don't immediately undergo a change, inter-penetration will result (figure 23).

How do we compute the change in velocity? Any force we might imagine acting at p, no matter how strong, would require at least a small amount of time to completely halt the relative motion between the bodies. (No matter how strong your car brakes are, you still need to apply them *before* you hit the brick wall. If you wait until you've contacted the wall, it's too late...) Since we want bodies to change their velocity instantly though, we postulate a new quantity J called an *impulse*. An impulse is a vector quantity, just like a force, but it has the units of momentum. Applying an impulse produces an instantaneous change in the velocity of a body. To determine the effects of a given impulse J, we imagine a large force F that acts for a small time interval  $\Delta t$ . If we let F go to infinity and  $\Delta t$  go to zero in such a way that

$$F\Delta t = J \tag{8-4}$$

then we can derive the effect of J on a body's velocity by considering how the velocity would change if we let the force F act on it for  $\Delta t$  time.

For example, if we apply an impulse J to a rigid body with mass M, then the change in linear

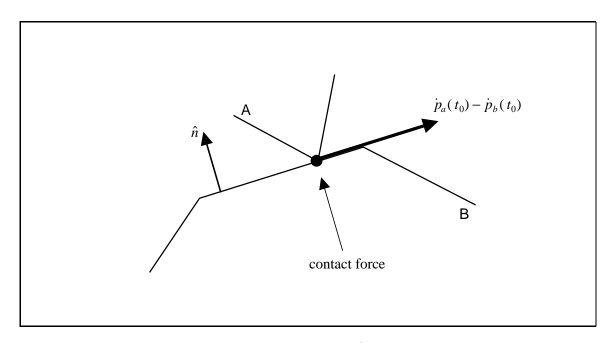


Figure 22: The vector  $\dot{p}_a(t_0) - \dot{p}_b(t_0)$  is perpendicular to  $\hat{n}(t_0)$ ; the bodies are in resting contact. A contact force may be necessary to prevent bodies from accelerating towards each other.

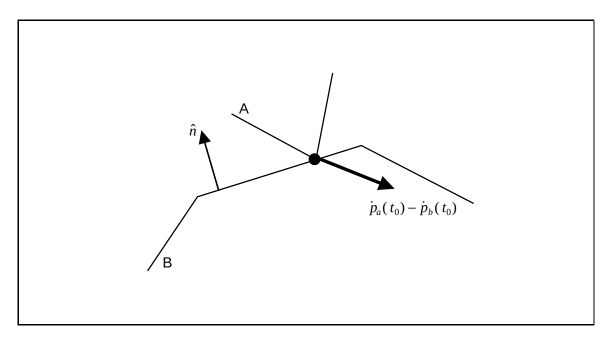


Figure 23: Colliding contact. The relative velocity  $\dot{p}_a(t_0) - \dot{p}_b(t_0)$  is directed inwards, opposite  $\hat{n}(t_0)$ . Unless the relative velocity is abruptly changed, inter-penetration will occur immediately after time  $t_0$ .

velocity  $\Delta v$  is simply

$$\Delta v = \frac{J}{M}.\tag{8-5}$$

Equivalently, the change in linear momentum  $\Delta P$  is simply  $\Delta P = J$ . If the impulse acts at the point p, then just as a force produces a torque, J produces an impulsive torque of

$$\tau_{impulse} = (p - x(t)) \times J. \tag{8-6}$$

As one would imagine, the impulsive torque  $\tau_{impulse}$  also gives rise to a change in angular momentum  $\Delta L$  of  $\Delta L = \tau_{impulse}$ . The change in angular velocity is simply  $I^{-1}(t_0)\tau_{impulse}$ , assuming the impulse was applied at time  $t_0$ .

When two bodies collide, we will apply an impulse between them to change their velocity. For frictionless bodies, the direction of the impulse will be in the normal direction,  $\hat{n}(t_0)$ . Thus, we can write the impulse J as

$$J = j\hat{n}(t_0) \tag{8-7}$$

where j is an (as yet) undetermined scalar that gives the magnitude of the impulse. We'll adopt the convention that the impulse J acts positively on body A, that is, A is subject to an impulse of  $+j\hat{n}(t_0)$ , while body B is subject to an equal but opposite impulse  $-j\hat{n}(t_0)$  (figure 24). We compute j by using an empirical law for collisions. Let's let  $\dot{p}_a^-(t_0)$  denote the velocity of the contact vertex of A prior to the impulse being applied, and let  $\dot{p}_a^+(t_0)$  denote the velocity after we apply the impulse J. Let  $\dot{p}_b^-(t_0)$  and  $\dot{p}_b^+(t_0)$  be defined similarly. Using this notation, the initial relative velocity in the normal direction is

$$v_{rel}^- = \hat{n}(t_0) \cdot (\dot{p}_a^-(t_0) - \dot{p}_b^-(t_0));$$
 (8–8)

after the application of the impulse,

$$v_{rel}^+ = \hat{n}(t_0) \cdot (\dot{p}_a^+(t_0) - \dot{p}_b^+(t_0)).$$
 (8–9)

The empirical law for frictionless collisions says simply that

$$v_{rel}^+ = -\epsilon v_{rel}^-. \tag{8-10}$$

The quantity  $\epsilon$  is called the *coefficient of restitution* and must satisfy  $0 \le \epsilon \le 1$ . If  $\epsilon = 1$ , then  $v_{rel}^+ = -v_{rel}^-$ , and the collision is perfectly "bouncy"; in particular, no kinetic energy is lost. At the other end of the spectrum,  $\epsilon = 0$  results in  $v_{rel}^+ = 0$ , and a maximum of kinetic energy is lost. After this sort of collision, the two bodies will be in resting contact at the contact point p (figure 25).

Calculating the magnitude j of the impulse  $J=j\hat{n}(t_0)$  is fairly simple, although the equations are a bit tedious to work through. Let's define the displacements  $r_a$  and  $r_b$  as  $p-x_a(t_0)$ , and  $p-x_b(t_0)$ . If we let  $v_a^-(t_0)$  and  $\omega_a^-(t_0)$  be the pre-impulse velocities of body A, and  $v_a^+(t_0)$  and  $\omega_a^+(t_0)$  be the post-impulse velocities, we can write

$$\dot{p}_a^+(t_0) = v_a^+(t_0) + \omega_a^+(t_0) \times r_a \tag{8-11}$$

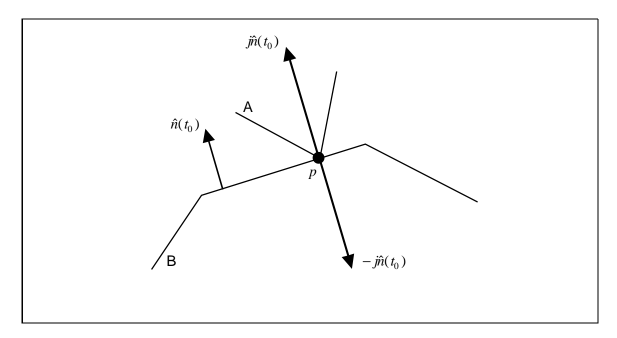


Figure 24: The impulse between two bodies at a contact point. An impulse of  $j\hat{n}(t_0)$  acts on A, while an impulse of  $-j\hat{n}(t_0)$  acts on B.

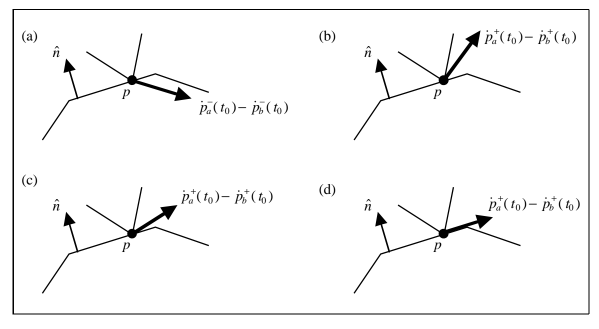


Figure 25: (a) The relative velocity before application of the impulse. (b) The component of the relative velocity in the  $\hat{n}(t_0)$  direction is reversed for an  $\epsilon=1$  collision. The relative velocity perpendicular to  $\hat{n}(t_0)$  remains the same. (c) A collision with  $0<\epsilon<1$ . The bodies bounce away in the  $\hat{n}(t_0)$  direction with less speed than they approached. (d) A collision with  $\epsilon=0$ . The bodies do not bounce away from each other, but the relative velocity perpendicular to  $\hat{n}(t_0)$  is unaffected by the collision.

along with

$$v_a^+(t_0) = v_a^-(t_0) + \frac{j\hat{n}(t_0)}{M_a}$$
 and  $\omega_a^+(t_0) = \omega_a^-(t_0) + I_a^{-1}(t_0) \left(r_a \times j\hat{n}(t_0)\right)$  (8-12)

where  $M_a$  is the mass of body A, and  $I_a(t_0)$  is its inertia tensor. Combining the two previous equations yields

$$\dot{p}_{a}^{+}(t_{0}) = \left(v_{a}^{-}(t_{0}) + \frac{j\hat{n}(t_{0})}{M_{a}}\right) + \left(\omega_{a}^{-}(t_{0}) + I_{a}^{-1}(t_{0})\left(r_{a} \times j\hat{n}(t_{0})\right)\right) \times r_{a}$$

$$= v_{a}^{-}(t_{0}) + \omega_{a}^{-}(t_{0}) \times r_{a} + \left(\frac{j\hat{n}(t_{0})}{M_{a}}\right) + \left(I_{a}^{-1}(t_{0})\left(r_{a} \times j\hat{n}(t_{0})\right)\right) \times r_{a}$$

$$= \dot{p}_{a}^{-} + j\left(\frac{\hat{n}(t_{0})}{M_{a}} + I_{a}^{-1}(t_{0})\left(r_{a} \times \hat{n}(t_{0})\right)\right) \times r_{a}.$$
(8-13)

It is important to note the form of  $\dot{p}_a^+(t_0)$ : it is a simple linear function of j. For body B, an opposite impulse  $-j\hat{n}(t_0)$  acts, yielding

$$\dot{p}_b^+(t_0) = \dot{p}_b^- - j \left( \frac{\hat{n}(t_0)}{M_b} + I_b^{-1}(t_0) \left( r_b \times \hat{n}(t_0) \right) \right) \times r_b.$$
 (8-14)

This yields

$$\dot{p}_{a}^{+}(t_{0}) - \dot{p}_{b}^{+} = \left(\dot{p}_{a}^{+}(t_{0}) - \dot{p}_{b}^{+}\right) + j\left(\frac{\hat{n}(t_{0})}{M_{a}} + \frac{\hat{n}(t_{0})}{M_{b}} + \left(I_{a}^{-1}(t_{0})\left(r_{a} \times \hat{n}(t_{0})\right)\right) \times r_{a} + \left(I_{b}^{-1}(t_{0})\left(r_{b} \times \hat{n}(t_{0})\right)\right) \times r_{b}\right).$$
(8-15)

To calculate  $v_{rel}^+$ , we dot this expression with  $\hat{n}(t_0)$ . Since  $\hat{n}(t_0)$  is of unit length,  $\hat{n}(t_0) \cdot \hat{n}(t_0) = 1$ , and we obtain

$$v_{rel}^{+} = \hat{n}(t_0) \cdot (\dot{p}_a^{+}(t_0) - \dot{p}_b^{+})$$

$$= \hat{n}(t_0) \cdot (\dot{p}_a^{-}(t_0) - \dot{p}_b^{-}) + j \left( \frac{1}{M_a} + \frac{1}{M_b} + \frac{1}{M_b} \right)$$

$$\hat{n}(t_0) \cdot \left( I_a^{-1}(t_0) \left( r_a \times \hat{n}(t_0) \right) \right) \times r_a + \hat{n}(t_0) \cdot \left( I_b^{-1}(t_0) \left( r_b \times \hat{n}(t_0) \right) \right) \times r_b$$

$$= v_{rel}^{-} + j \left( \frac{1}{M_a} + \frac{1}{M_b} + \frac{1}{M_b} \right)$$

$$\hat{n}(t_0) \cdot \left( I_a^{-1}(t_0) \left( r_a \times \hat{n}(t_0) \right) \right) \times r_a + \hat{n}(t_0) \cdot \left( I_b^{-1}(t_0) \left( r_b \times \hat{n}(t_0) \right) \right) \times r_b$$
(8-16)

By expressing  $v_{rel}^+$  in terms of j and  $v_{rel}^-$ , we can compute j according to equation (8–10). If we substitute equation (8–16) into equation (8–10), we get

$$v_{rel}^{-} + j \left( \frac{1}{M_a} + \frac{1}{M_b} + \hat{n}(t_0) \cdot \left( I_a^{-1}(t_0) \left( r_a \times \hat{n}(t_0) \right) \right) \times r_a + \hat{n}(t_0) \cdot \left( I_b^{-1}(t_0) \left( r_b \times \hat{n}(t_0) \right) \right) \times r_b \right) = -\epsilon v_{rel}^{-}.$$
(8-17)

Finally, solving for j,

```
j = \frac{-(1+\epsilon)v_{rel}^{-}}{\frac{1}{M_a} + \frac{1}{M_b} + \hat{n}(t_0) \cdot \left(I_a^{-1}(t_0)\left(r_a \times \hat{n}(t_0)\right)\right) \times r_a + \hat{n}(t_0) \cdot \left(I_b^{-1}(t_0)\left(r_b \times \hat{n}(t_0)\right)\right) \times r_b}. (8–18)
```

Let's consider some actual code (written for clarity, not speed). First, we determine if two bodies are in colliding contact.

```
/*
 * Operators: if 'x' and 'y' are triples,
 * assume that 'x ^ y' is their cross product,
 * and 'x * y' is their dot product.
 * /
/* Return the velocity of a point on a rigid body */
triple pt_velocity(Body *body, triple p)
    return body->v + (body->omega ^ (p - body->x));
}
/*
 * Return true if bodies are in colliding contact. The
 * parameter 'THRESHOLD' is a small numerical tolerance
 * used for deciding if bodies are colliding.
 * /
bool colliding(Contact *c)
    triple padot = pt_velocity(c->a, p), /* \dot{p}_a^-(t_0) */ pbdot = pt_velocity(c->b, p); /* \dot{p}_b^-(t_0) */
    double vrel = c->n * (padot - pbdot); /* v_{rel}^- */
    if(vrel > THRESHOLD) /* moving away */
        return false;
    if(vrel > -THRESHOLD) /* resting contact */
        return false;
                              /* vrel < -THRESHOLD */</pre>
    else
        return true;
}
```

Next, we'll loop through all the contact points until all the collisions are resolved, and actually compute and apply an impulse.

```
void collision(Contact *c, double epsilon)  \{ \\  \text{triple padot} = \text{pt\_velocity(c->a, c->p), /* } \dot{p}_a^-(t_0) \text{ */} \\  \text{pbdot} = \text{pt\_velocity(c->b, c->p), /* } \dot{p}_b^-(t_0) \text{ */} \\  n = \text{c->n,} \text{ /* } \hat{n}(t_0) \text{ */}
```

```
/* r_a */
            ra = p - c->a->x,
                                               /* r_b */
            rb = p - c -> b -> x;
                                              /* v<sub>rel</sub> */
    double vrel = n * (padot - pbdot),
            numerator = -(1 + epsilon) * vrel;
    /* We'll calculate the denominator in four parts */
    double term1 = 1 / c \rightarrow a \rightarrow mass,
            term2 = 1 / c->b->mass,
            term3 = n * ((c->a->Iinv * (ra ^ n)) ^ ra),
            term4 = n * ((c->b->Iinv * (rb ^ n)) ^ rb);
    /* Compute the impulse magnitude */
    double j = numerator / (term1 + term2 + term3 + term4);
    triple force = j * n;
    /* Apply the impulse to the bodies */
    c->a->P += force;
    c->b->P -= force;
    c->a->L += ra ^ force;
    c->b->L -= rb ^ force;
    /* recompute auxiliary variables */
    c->a->v = c->a->P / c->a->mass;
    c->b->v = c->b->P / c->b->mass;
    c->a->omega = c->a->Iinv * c->a->L;
    c->b->omega = c->b->Iinv * c->b->L;
}
void FindAllCollisions(Contact contacts[], int ncontacts)
    bool had_collision;
    double epsilon = .5;
    do {
        had collision = false;
        for(int i = 0; i < ncontacts; i++)</pre>
            if(colliding(&contacts[i]))
                collision(&contacts[i], epsilon);
                had_collision = true;
                /* Tell the solver we had a collision */
                ode_discontinuous();
```

```
} while(had_collision == true);
}
```

Note several things. First,  $\epsilon = .5$  was chosen arbitrarily. In a real implementation, we'd allow the user to use different values of  $\epsilon$  depending on which two bodies were colliding. Also, every time we find a collision, we have to rescan the list of contacts, since bodies that were at rest may no longer be so, and new collisions may develop. If there are initially several collisions to be resolved (such as a cube dropped flat onto a plane, with all four vertices colliding at once), the order of the contact list may have an effect on the simulation. There is a way to compute impulses at more than one contact point at a time, but it more complicated, and is based on the concepts used for resting contact in the next section. For further information, see Baraff[1].

Incidentally, if you want to have certain bodies that are "fixed", and cannot be moved (such as floors, or walls), you can use the following trick: for such bodies, let  $\frac{1}{\text{mass}}$  be zero; also let the inverse inertia tensor also be the  $3 \times 3$  zero matrix. You can either special-case the code to check if a body is supposed to be fixed, or you can recode the definition of RigidBody to have the variable invmass instead of mass. For ordinary bodies, invmass is the inverse of the mass, while for fixed bodies, invmass is zero. The same goes for the inertia tensor. (Note that nowhere in any of the dynamics computations (including the next section) is the mass or inertia tensor ever used; only their inverses are used, so you won't have to worry about dividing by zero.) The same trick can be used in the next section on resting contact to simulate bodies that can support any amount of weight without moving.

# 9 Resting Contact

The case of resting contact, when bodies are neither colliding nor separating at a contact point, is the last (and hardest) dynamics problem we'll tackle in these notes. To implement what's in this section, you'll have to obtain a fairly sophisticated piece of numerical software, which we'll describe below.

At this point, let's assume we have a configuration with n contact points. At each contact point, bodies are in resting contact, that is, the relative velocity  $v_{rel}$ , from section 8, is zero (to within the numerical tolerance THRESHOLD). We can say that this is so, because colliding contact is eliminated by the routine FindAllCollisions(), and any contact points with  $v_{rel}$  larger than THRESHOLD can be safely ignored, since the bodies are separating there.

As was the case for colliding contact, at each contact point, we have a contact force that acts normal to the contact surface. For the case of colliding contact, we had an impulse  $j\hat{n}(t_0)$  where j was an unknown scalar. For resting contact, at each contact point there is some force  $f_i\hat{n}_i(t_0)$ , where  $f_i$  is an unknown scalar, and  $\hat{n}_i(t_0)$  is the normal at the ith contact point (figure 26). Our goal is to determine what each  $f_i$  is. In computing the  $f_i$ 's, they must all be determined at the same time, since the force at the ith contact point may influence one or both of the bodies of the j contact point. In section 8, we wrote how the velocity of the contact points  $p_a(t_0)$  and  $p_b(t_0)$  changed with respect to j. We'll do the same thing here, but now we'll have to describe how the acceleration of  $p_a(t_0)$  and  $p_b(t_0)$  depends on each  $f_i$ .

For colliding contact, we had an empirical law which related the impulse strength j to the relative velocity and a coefficient of restitution. For resting contact, we compute the  $f_i$ 's subject to not one, but three conditions. First, the contact forces must prevent inter-penetration; that is, the contact

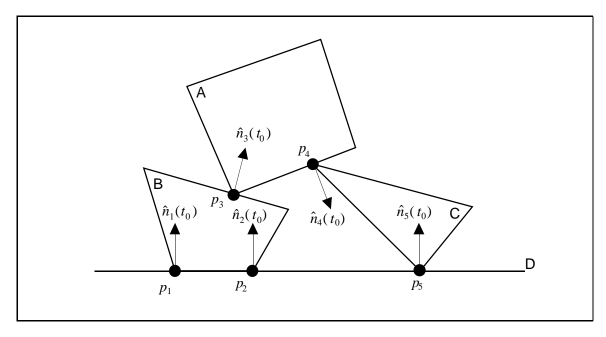


Figure 26: Resting contact. This configuration has five contact points; a contact force acts between pairs of bodies at each contact point.

forces must be strong enough to prevent two bodies in contact from being pushed "towards" one another. Second, we want our contact forces to be repulsive; contact forces can push bodies apart, but can never act like "glue" and hold bodies together. Last, we require that the force at a contact point become zero if the bodies begin to separate. For example, if a block is resting on a table, some force may act at each of the contact points to prevent the block from accelerating downwards in response to the pull of gravity. However, if a very strong wind were to blow the brick upwards, the contact forces on the brick would have to become zero at the instant that the wind accelerated the brick off the table.

Let's deal with the first condition: preventing inter-penetration. For each contact point i, we construct an expression  $d_i(t)$  which describes the separation distance between the two bodies near the contact point at time t. Positive distance indicates the bodies have broken contact, and have separated at the ith contact point, while negative distance indicates inter-penetration. Since the bodies are in contact at the present time  $t_0$ , we will have  $d_i(t_0) = 0$  (within numerical tolerances) for each contact point. Our goal is to make sure that the contact forces maintain  $d_i(t) \ge 0$  for each contact point at future times  $t > t_0$ .

For vertex/face contacts, we can immediately construct a very simple function for  $d_i(t)$ . If  $p_a(t)$  and  $p_b(t)$  are the contact points of the *i*th contact, between bodies A and B, than the distance between the vertex and the face at future times  $t \ge t_0$  is given by

$$d_i(t) = \hat{n}_i(t) \cdot (p_a(t) - p_b(t)). \tag{9-1}$$

At time t, the function d(t) measures the separation between A and B near  $p_a(t)$ . If  $d_i(t)$  is zero, then the bodies are in contact at the ith contact point. If  $d_i(t) > 0$ , then the bodies have lost contact at the ith contact point. However, if  $d_i(t) < 0$ , then the bodies have inter-penetrated, which is what we need to avoid (figure 27). The same function can also be used for edge/edge contacts; since  $\hat{n}_i(t)$ 

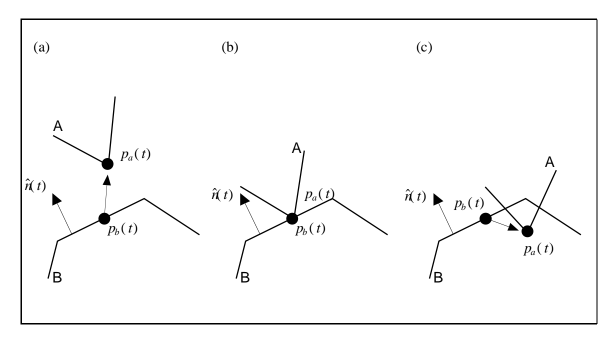


Figure 27: (a) The displacement  $p_a(t) - p_b(t)$ , indicated by an arrow, points in the same direction as  $\hat{n}(t)$ . Thus, the distance function d(t) would be positive. (b) The distance function d(t) is zero. (c) The displacement  $p_a(t) - p_b(t)$  points in the opposite direction as  $\hat{n}(t)$ . The distance function d(t) is negative, indicating inter-penetration.

points outwards from B towards A (by convention),  $\hat{n}_i(t) \cdot (p_a(t) - p_b(t))$  will be positive if the two contacting edges move so as to separate the bodies.

Since  $d_i(t_0) = 0$ , we have to keep  $d_i(t_0)$  from decreasing at time  $t_0$ ; that is, we have to have  $\dot{d}_i(t_0) \ge 0$ . What is  $\dot{d}_i(t_0)$ ? Differentiating,

$$\dot{d}_i(t) = \dot{\hat{n}}_i(t) \cdot (p_a(t) - p_b(t)) + \hat{n}_i(t) \cdot (\dot{p}_a(t) - \dot{p}_b(t)). \tag{9-2}$$

Since  $d_i(t)$  describes the separation distance,  $\dot{d}_i(t)$  will describe the separation velocity at time t. However, at time  $t_0$ ,  $p_a(t_0) = p_b(t_0)$ , which means that  $\dot{d}_i(t_0) = \hat{n}_i(t_0) \cdot (\dot{p}_a(t_0) - \dot{p}_b(t_0))$ . This should look familiar: its  $v_{rel}$  from the previous section! The function  $\dot{d}_i(t_0)$  is a measure of how the bodies are separating, and for resting contact, we know that  $\dot{d}(t_0)$  is zero, because the bodies are neither moving towards nor away from each other at a contact point.

At this point then, we have  $d_i(t_0) = \dot{d}_i(t_0) = 0$ . Now we'll look at  $\ddot{d}_i(t_0)$ . If we differentiate equation (9–2), we get

$$\begin{split} \ddot{d_i}(t) &= \left( \ddot{\hat{n}}_i(t) \cdot (p_a(t) - p_b(t)) + \dot{\hat{n}}_i(t) \cdot (\dot{p}_a(t) - \dot{p}_b(t)) \right) + \\ &\left( \dot{\hat{n}}_i(t) \cdot (\dot{p}_a(t) - \dot{p}_b(t)) + \hat{n}_i(t) \cdot (\ddot{p}_a(t) - \ddot{p}_b(t)) \right) \\ &= \ddot{\hat{n}}_i(t) \cdot (p_a(t) - p_b(t)) + 2\dot{\hat{n}}_i(t) \cdot (\dot{p}_a(t) - \dot{p}_b(t)) + \hat{n}_i(t) \cdot (\ddot{p}_a(t) - \ddot{p}_b(t)). \end{split}$$
(9-3)

Since  $p_a(t_0) = p_b(t_0)$ , we can write  $\ddot{d}_i(t_0)$  as

$$\ddot{d}(t_0) = \hat{n}_i(t_0) \cdot (\ddot{p}_a(t_0) - \ddot{p}_b(t_0)) + 2\dot{\hat{n}}_i(t_0) \cdot (\dot{p}_a(t_0) - \dot{p}_b(t_0)). \tag{9-4}$$

The quantity  $\ddot{d}_i(t_0)$  measures how the two bodies are accelerating towards each other at the contact point p. If  $\ddot{d}_i(t_0) > 0$ , the the bodies have an acceleration away from each other, and contact will break immediately after  $t_0$ . If  $\ddot{d}_i(t_0) = 0$ , then contact remains. The case  $\ddot{d}_i(t_0) < 0$  must be avoided, for this indicates the bodies are accelerating towards each other. Note that if  $\hat{n}_i(t_0)$  is a constant (if body B is fixed), then  $\dot{n}_i(t_0)$  is zero, leading to further simplifications.

Thus, we satisfy our first condition for contact forces by writing the constraint

$$\ddot{d}_i(t_0) \ge 0 \tag{9-5}$$

for each contact point. Since the acceleration  $\ddot{d}_i(t_0)$  depends on the contact forces, this is really a constraint on the contact forces.

Let's turn our attention to the second and third constraints. Since contact forces must always be repulsive, each contact force must act outward. This means that each  $f_i$  must be positive, since a force of  $f_i\hat{n}_i(t_0)$  acts on body A, and  $\hat{n}_i(t_0)$  is the outwards pointing normal of B. Thus, we need

$$f_i \ge 0 \tag{9-6}$$

for each contact point. The third constraint is expressed simply in terms of  $f_i$  and  $\ddot{d}_i(t_0)$ . Since the contact force  $f_i\hat{n}_i(t_0)$  must become zero if contact is breaking at the *i*th contact, this says that  $f_i$  must be zero if contact is breaking. We can express this constraint by writing

$$f_i \ddot{d}_i(t_0) = 0;$$
 (9–7)

if contact is breaking,  $\ddot{d}_i(t_0) > 0$  and equation (9–7) is satisfied by requiring  $f_i = 0$ . If contact is not breaking, then  $\ddot{d}_i(t_0) = 0$ , and equation (9–7) is satisfied regardless of  $f_i$ .

In order to actually find  $f_i$ 's which satisfy equations (9–5), (9–6), and (9–7), we need to express each  $\ddot{d}_i(t_0)$  as a function of the unknown  $f_i$ 's. It will turn out that we will be able to write each  $\ddot{d}_i(t_0)$  in the form

$$\ddot{d}_i(t_0) = a_{i1}f_1 + a_{i2}f_2 + \dots + a_{in}f_n + b_i.$$
(9-8)

In matrix parlance, this means we will be able to write

$$\begin{pmatrix} \ddot{d}_1(t_0) \\ \vdots \\ \ddot{d}_n(t_0) \end{pmatrix} = \mathbf{A} \begin{pmatrix} f_1 \\ \vdots \\ f_n \end{pmatrix} + \begin{pmatrix} b_i \\ \vdots \\ b_n \end{pmatrix}$$
(9-9)

where **A** is the  $n \times n$  matrix of the  $a_{ij}$  coefficients of equation (9–8). Although the code needed to calculate the  $a_{ij}$ 's and the  $b_i$ 's is not too complicated, working out the derivations on which the code is based is somewhat tedious. The derivations are worked out in appendix D, along with code to compute the matrix of  $a_{ij}$ 's and  $b_i$ 's.

Appendix D gives an implementation of the routines

where the types bigmatrix and vector represent matrices and vectors of arbitrary size. The first routine computes the  $a_{ii}$ 's, while the second routine computes the  $b_i$ 's.

Once we've computed all this, we can think about solving equations (9–5), (9–6), and (9–7). This system of equations forms what is called a *quadratic program* (QP); that is,  $f_i$ 's that satisfy these three equations are found by an algorithm called quadratic programming. Not all quadratic programs can be solved efficiently, but because our contact forces are all normal to the contact surfaces (that is, they do not involve friction), it turns out that our QP can always be solved efficiently. One interesting thing to note is that QP codes can easily handle the case  $\ddot{d}_i(t_0) = 0$  instead of  $\ddot{d}_i(t_0) \geq 0$ . We use  $\ddot{d}_i(t_0) = 0$  (and also drop the constraint  $f_i \geq 0$ ) if we wish to constrain two bodies to never separate at a contact point. This enables us to implement hinges, and pin-joints, as well as non-penetration constraints during simulation.

Quadratic programming codes aren't terribly common though; certainly, they are not nearly as common as linear equation codes, and are much harder to implement. The quadratic programming routines used by the author were obtained from the Department of Operations Research at Stanford University. See Gill *et al.*[7, 8, 9] for further details. More recently, we have been using code described by Baraff[3] to solve the quadratic programs. If you are determined to really implement this, we suggest a thorough study of this paper (excepting for the section on contact with friction).

At any rate, let's assume that you've got a working QP solver at your disposal. We'll assume that you pass the matrix A, and the vector of  $b_i$ 's to the QP solver, and you get back the vector of  $f_i$ 's. Let's pretend the interface is

```
void qp_solve(bigmatrix &a, vector &b, vector &f);
```

Let's see how to compute all the resting contact forces. The following routine is presumably called from ComputeForceAndTorque(), after FindAllCollisions() has been called.

```
bigmatrix amat = new bigmatrix(ncontacts, ncontacts);
             bvec = new vector(ncontacts),
   vector
              fvec = new vector(ncontacts);
    /* Compute a_{ij} and b_i coefficients */
    compute_a_matrix(contacts, ncontacts, amat);
    compute_b_vector(contacts, ncontacts, bvec);
    /* Solve for f_i's */
   qp_solve(amat, bmat, fvec);
    /* Now add the resting contact forces we just computed into
       the 'force' and 'torque' field of each rigid body. */
   for(int i = 0; i < ncontacts; i++)</pre>
       double f = fvec[i];
                                               /* f_i */
                 n = contacts[i]->n;
                                              /* \hat{n}_i(t_0) * /
        triple
                                               /* body A */
       RigidBody *A = contacts[i]->a,
                   *B = contacts[i]->b;
                                             /* body B */
        /* apply the force `f n' positively to A... */
       A->force += f * n;
       A->torque += (contacts[i].p - A->x) * (f*n);
        /* and negatively to B */
       B->force -= f * n;
       B->torque -= (contacts[i].p - B->x) * (f*n);
    }
}
```

That's pretty much it! Now that the resting forces have been computed and combined with the external forces, we return control to the ODE solver, and each body goes merrily along its way, in a physically correct manner, without inter-penetration.

### **Appendix A** Motion Equation Derivations

In this appendix, we'll fill in some of the missing details from section 2, with regards to the equations  $\dot{P}(t) = F(t)$ ,  $\dot{L}(t) = \tau(t)$ , and  $L(t) = I(t)\omega(t)$ . The derivation method used here is somewhat nonstandard, and was proposed by Andy Witkin. The derivation in this appendix is (we feel) much shorter and considerably more elegant than the one found in traditional sources such as Goldstein[10].

We've described the external force acting on a rigid body in terms of forces  $F_i(t)$ , where  $F_i(t)$  is the external force acting on the *i*th particle. However, for a rigid body to maintain its shape, there must be some "internal" constraint forces that act between particles in the same body. We will make the assumption that these constraint forces act passively on the system and do not perform any net work. Let  $F_{ci}(t)$  denote the net internal constraint force acting on the *i*th particle. The work performed by  $F_{ci}$  on the *i*th particle from time  $t_0$  to  $t_1$  is

$$\int_{t_0}^{t_1} F_{ci}(t) \cdot \dot{r}_i(t) dt$$

where  $\dot{r}_i(t)$  is the velocity of the *i*th particle. The net work over all the particles is the sum

$$\sum_{i} \int_{t_0}^{t_1} F_{ci}(t) \cdot \dot{r}_i(t) dt = \int_{t_0}^{t_1} \sum_{i} F_{ci}(t) \cdot \dot{r}_i(t) dt,$$

which must be zero for any interval  $t_0$  to  $t_1$ . This means that the integrand

$$\sum_{i} F_{ci}(t) \cdot \dot{r}_{i}(t) \tag{A-1}$$

is itself always zero for any time t. (Henceforth we'll just write these expressions as  $\sum F_{ci} \cdot \dot{r}_i = 0$ .) We can use this fact to eliminate any mention of the constraint forces  $F_{ci}$  from our derivations.

First, some quick notes about the "\*" operator defined in section 2.3: since  $a^*b = a \times b$ , and  $a \times b = -b \times a$ , we get

$$-a^*b = b \times a = b^*a. \tag{A-2}$$

Since  $a^*$  is an antisymmetric matrix,

$$(a^*)^T = -a^*. (A-3)$$

Finally, since the "\*" operator is a linear operator,

$$(\dot{a})^* = (\dot{a^*}) = \frac{d}{dt}(a^*)$$
 (A-4)

and for a set of vectors  $a_i$ 

$$\sum a_i^* = \left(\sum a_i\right)^*. \tag{A-5}$$

Recall that we can write the velocity  $\dot{r}_i$  as  $\dot{r}_i = v + \omega \times (r_i - x)$  where  $r_i$  is the particle's location, x is the position of the center of mass, and v and  $\omega$  are linear and angular velocity. Letting  $r'_i = r_i - x$  and using the "\*" notation,

$$\dot{r}_i = v + \omega^* r_i' = v - r_i'^* \omega. \tag{A-6}$$

Substituting this into  $\sum F_{ci} \cdot \dot{r}_i$ , which is always zero, yields

$$\sum F_{ci} \cdot (v - r_i^{\prime *} \omega) = 0. \tag{A-7}$$

$$\sum -F_{ci}^{T}r_{i}^{\prime*}\omega = \left(\sum -F_{ci}^{T}r_{i}^{\prime*}\right)\omega = 0 \tag{A-8}$$

for any  $\omega$ , so  $\sum -F_{ci}^T r_i^{\prime*} = \mathbf{0}^T$ . Transposing, we have

$$\sum -(r_i^{\prime*})^T F_{ci} = \sum (r_i^{\prime})^* F_{ci} = \sum r_i^{\prime} \times F_{ci} = \mathbf{0}$$
 (A-9)

which means that the internal forces produce no net torque.

We can use the above to derive the rigid body equations of motion. The net force on each particle is the sum of the internal constraint force  $F_{ci}$  and the external force  $F_i$ . The acceleration  $\ddot{r}_i$  of the *i*th particle is

$$\ddot{r}_i = \frac{d}{dt}\dot{r}_i = \frac{d}{dt}(v - r_i^{\prime*}\omega) = \dot{v} - \dot{r}_i^{\prime*}\omega - r_i^{\prime*}\dot{\omega}. \tag{A-10}$$

Since each individual particle must obey Newton's law f = ma, or equivalently ma - f = 0, we have

$$m_i \ddot{r}_i - F_i - F_{ci} = m_i (\dot{v} - \dot{r}_i^{\prime *} \omega - r_i^{\prime *} \dot{\omega}) - F_i - F_{ci} = \mathbf{0}$$
 (A-11)

for each particle.

To derive  $\dot{P} = F = \sum F_i$ , we sum equation (A–11) over all the particles. We obtain

$$\sum m_i (\dot{v} - \dot{r}_i^{\prime *} \omega - r_i^{\prime *} \dot{\omega}) - F_i - F_{ci} = \mathbf{0}.$$
 (A-12)

Breaking the large sum into smaller ones,

$$\sum m_{i}(\dot{v} - \dot{r}_{i}^{\prime*}\omega - r_{i}^{\prime*}\dot{\omega}) - F_{i} - F_{ci} =$$

$$\sum m_{i}\dot{v} - \sum m_{i}\dot{r}_{i}^{\prime*}\omega - \sum m_{i}r_{i}^{\prime*}\dot{\omega} - \sum F_{i} - \sum F_{ci} =$$

$$\sum m_{i}\dot{v} - \left(\sum m_{i}\dot{r}_{i}^{\prime}\right)^{*}\omega - \left(\sum m_{i}r_{i}^{\prime}\right)^{*}\dot{\omega} - \sum F_{i} - \sum F_{ci} =$$

$$\sum m_{i}\dot{v} - \left(\frac{d}{dt}\sum m_{i}r_{i}^{\prime}\right)^{*}\omega - \left(\sum m_{i}r_{i}^{\prime}\right)^{*}\dot{\omega} - \sum F_{i} - \sum F_{ci} = \mathbf{0}.$$
(A-13)

Since we are in a center-of-mass coordinate system, equation (2–20) from section 2.6 tells us that  $\sum m_i r_i' = \mathbf{0}$ , which also means that  $\frac{d}{dt} \sum m_i r_i' = \mathbf{0}$ . Removing terms with  $\sum m_i r_i'$ , and the term  $\sum F_{ci}$  from the above equation yields

$$\sum m_i \dot{v} - \sum F_i = \mathbf{0} \tag{A-14}$$

or simply  $M\dot{v} = \dot{P} = \sum F_i$  as advertised.

To obtain  $\dot{L} = \tau = \sum r'_i \times F_i$ , we again start with equation (A–11). Multiplying both sides by  $r'^*_i$  yields

$$r_i^{\prime *} m_i (\dot{v} - \dot{r}_i^{\prime *} \omega - r_i^{\prime *} \dot{\omega}) - r_i^{\prime *} F_i - r_i^{\prime *} F_{ci} = r_i^{\prime *} \mathbf{0} = \mathbf{0}.$$
 (A-15)

Summing over all the particles, we obtain

$$\sum r_i^{\prime *} m_i \dot{v} - \sum r_i^{\prime *} m_i \dot{r}_i^{\prime *} \omega - \sum r_i^{\prime *} m_i r_i^{\prime *} \dot{\omega} - \sum r_i^{\prime *} F_i - \sum r_i^{\prime *} F_{ci} = \mathbf{0}.$$
 (A-16)

Since  $\sum r_i^{\prime *} F_{ci} = \mathbf{0}$ , we can rearrange this to obtain

$$\left(\sum m_i r_i'\right)^* \dot{v} - \left(\sum m_i r_i'^* \dot{r}_i'^*\right) \omega - \left(\sum m_i r_i'^* r_i'^*\right) \dot{\omega} - \sum r_i'^* F_i = \mathbf{0}. \tag{A-17}$$

Using  $\sum m_i r'_i = \mathbf{0}$ , we are left with

$$-\left(\sum m_{i}r_{i}^{\prime*}\dot{r}_{i}^{\prime*}\right)\omega - \left(\sum m_{i}r_{i}^{\prime*}r_{i}^{\prime*}\right)\dot{\omega} - \sum r_{i}^{\prime*}F_{i} - = \mathbf{0}$$
 (A-18)

or, recognizing that  $\sum r_i^{\prime *} F_i = \sum r_i^{\prime} \times F_i = \tau$ ,

$$-\left(\sum m_i r_i'^* \dot{r}_i'^*\right) \omega - \left(\sum m_i r_i'^* r_i'^*\right) \dot{\omega} = \tau. \tag{A-19}$$

We're almost done now: if we refer back to the matrix defined by the "\*" notation, one can easily verify the relation that the matrix  $-a^*a^*$  is equivalent to the matrix  $(a^Ta)\mathbf{1} - aa^T$  where  $\mathbf{1}$  is the  $3 \times 3$  identity matrix. (This relation is equivalent to the vector rule  $a \times (b \times c) = ba^Tc - ca^Tb$ .) Thus

$$\sum -m_i r_i'^* r_i'^* = \sum m_i ((r_i'^T r_i') \mathbf{1} - r_i' r_i'^T) = I(t).$$
 (A-20)

Substituting into equation (A–19), this yields

$$\left(\sum -m_i r_i^{\prime *} \dot{r}_i^{\prime *}\right) \omega + I(t) \dot{\omega} = \tau. \tag{A-21}$$

The above expression is almost acceptable, as it gives an expression for  $\dot{\omega}$  in terms of  $\tau$ , except that it requires us to evaluate the matrix  $\sum m_i r_i'^* \dot{r}_i^*$ , which is as expensive as computing the inertia tensor from scratch. We'll use one last trick here to clean things up. Since  $\dot{r}_i' = \omega \times r_i'$  and  $r_i'^* \omega = -\omega \times r_i'$ , we can write

$$\sum m_i \dot{r}_i'^* r_i'^* \omega = \sum m_i (\omega \times r_i')^* (-\omega \times r_i') = \sum -m_i (\omega \times r_i') \times (\omega \times r_i') = \mathbf{0}.$$
 (A-22)

Thus, we can add  $-\sum m_i \dot{r}_i^{\prime *} r_i^{\prime *} \omega = \mathbf{0}$  to equation (A–21) to obtain

$$\left(\sum -m_i r_i^{\prime *} \dot{r}_i^{\prime *} - m_i \dot{r}_i^{\prime *} r_i^{\prime *}\right) \omega + I(t) \dot{\omega} = \tau. \tag{A-23}$$

Finally, since

$$\dot{I}(t) = \frac{d}{dt} \sum -m_i r_i^{\prime *} r_i^{\prime *} = \sum -m_i r_i^{\prime *} \dot{r}_i^{\prime *} - m_i \dot{r}_i^{\prime *} r_i^{\prime *}$$
(A-24)

we have

$$\dot{I}(t)\omega + I(t)\dot{\omega} = \frac{d}{dt}(I(t)\omega) = \tau.$$
 (A-25)

Since  $L(t) = I(t)\omega(t)$ , this leaves us with the final result that

$$\dot{L}(t) = \tau. \tag{A-26}$$

# **Appendix B Quaternion Derivations**

A formula for  $\dot{q}(t)$  is derived as follows. Recall that the angular velocity  $\omega(t)$  indicates that the body is instantaneously rotating about the  $\omega(t)$  axis with magnitude  $|\omega(t)|$ . Suppose that a body were to rotate with a constant angular velocity  $\omega(t)$ . Then the rotation of the body after a period of time  $\Delta t$  is represented by the quaternion

$$\left[\cos\frac{|\omega(t)|\Delta t}{2}, \sin\frac{|\omega(t)|\Delta t}{2}\frac{\omega(t)}{|\omega(t)|}\right].$$

Let us compute  $\dot{q}(t)$  at some particular instant of time  $t_0$ . At times  $t_0 + \Delta t$  (for small  $\Delta t$ ), the orientation of the body is (to within first order) the result of first rotating by  $q(t_0)$  and then further rotating with velocity  $\omega(t_0)$  for  $\Delta t$  time. Combining the two rotations, we get

$$q(t_0 + \Delta t) = \left[\cos \frac{|\omega(t_0)|\Delta t}{2}, \sin \frac{|\omega(t_0)|\Delta t}{2} \frac{\omega(t_0)}{|\omega(t_0)|}\right] q(t_0).$$
 (B-1)

Making the substitution  $t = t_0 + \Delta t$ , we can express this as

$$q(t) = \left[\cos\frac{|\omega(t_0)|(t-t_0)}{2}, \sin\frac{|\omega(t_0)|(t-t_0)}{2} \frac{\omega(t_0)}{|\omega(t_0)|}\right] q(t_0).$$
 (B-2)

Let us differentiate q(t) at time  $t_0$ . First, since  $q(t_0)$  is a constant, let us differentiate

$$\left[\cos\frac{|\omega(t_0)|(t-t_0)}{2}, \sin\frac{|\omega(t_0)|(t-t_0)}{2} \frac{\omega(t_0)}{|\omega(t_0)|}\right].$$

At time  $t = t_0$ ,

$$\frac{d}{dt}\cos\frac{|\omega(t_0)|(t-t_0)}{2} = -\frac{|\omega(t_0)|}{2}\sin\frac{|\omega(t_0)|(t-t_0)}{2} 
= -\frac{|\omega(t_0)|}{2}\sin 0 = 0.$$
(B-3)

Similarly,

$$\frac{d}{dt}\sin\frac{|\omega(t_0)|(t-t_0)}{2} = \frac{|\omega(t_0)|}{2}\cos\frac{|\omega(t_0)|(t-t_0)}{2} 
= \frac{|\omega(t_0)|}{2}\cos 0 = \frac{|\omega(t_0)|}{2}.$$
(B-4)

Thus, at time  $t = t_0$ ,

$$\dot{q}(t) = \frac{d}{dt} \left( \left[ \cos \frac{|\omega(t_0)|(t-t_0)}{2}, \sin \frac{|\omega(t_0)|(t-t_0)}{2} \frac{\omega(t_0)}{|\omega(t_0)|} \right] q(t_0) \right) 
= \frac{d}{dt} \left( \left[ \cos \frac{|\omega(t_0)|(t-t_0)}{2}, \sin \frac{|\omega(t_0)|(t-t_0)}{2} \frac{\omega(t_0)}{|\omega(t_0)|} \right] \right) q(t_0) 
= \left[ 0, \frac{|\omega(t_0)|}{2} \frac{\omega(t_0)}{|\omega(t_0)|} \right] q(t_0) 
= \left[ 0, \frac{1}{2} \omega(t_0) \right] q(t_0) = \frac{1}{2} \left[ 0, \omega(t_0) \right] q(t_0).$$
(B-5)

The product  $[0, \omega(t_0)]q(t_0)$  is abbreviated to the form  $\omega(t_0)q(t_0)$ ; thus, the general expression for  $\dot{q}(t)$  is

$$\dot{q}(t) = \frac{1}{2}\omega(t)q(t). \tag{B-6}$$

### Appendix C Some Miscellaneous Formulas

### C.1 Kinetic Energy

The kinetic energy T of a rigid body is defined as

$$T = \sum_{i=1}^{n} \frac{1}{2} m_i \dot{r}_i^T \dot{r}_i. \tag{C-1}$$

Letting  $r'_i = r_i - x$ , we have  $\dot{r}_i = v(t) + r'^*_i \omega$ . Thus

$$T = \sum_{i=1}^{\infty} \frac{1}{2} m_{i} \dot{r}_{i}^{T} \dot{r}_{i}$$

$$= \sum_{i=1}^{\infty} \frac{1}{2} m_{i} (v + r_{i}^{\prime *} \omega)^{T} (v + r_{i}^{\prime *} \omega)$$

$$= \frac{1}{2} \sum_{i=1}^{\infty} m_{i} v^{T} v + \sum_{i=1}^{\infty} v^{T} m_{i} r_{i}^{\prime *} \omega + \frac{1}{2} \sum_{i=1}^{\infty} m_{i} (r_{i}^{\prime *} \omega)^{T} (r_{i}^{\prime *} \omega)$$

$$= \frac{1}{2} v^{T} \left(\sum_{i=1}^{\infty} m_{i}\right) v + v^{T} \left(\sum_{i=1}^{\infty} m_{i} r_{i}^{\prime}\right)^{*} \omega + \frac{1}{2} \omega^{T} \left(\sum_{i=1}^{\infty} m_{i} (r_{i}^{\prime *})^{T} r_{i}^{\prime *}\right) \omega.$$
(C-2)

Using  $\sum m_i r_i' = \mathbf{0}$  and  $(r_i'^*)^T = -r_i'^*$ , we have

$$T = \frac{1}{2}v^{T}Mv + \frac{1}{2}\omega^{T}\left(\sum -m_{i}r_{i}^{\prime*}r_{i}^{\prime*}\right)\omega = \frac{1}{2}(v^{T}Mv + \omega^{T}I\omega)$$
 (C-3)

since  $I = \sum -m_i r_i^{\prime *} r_i^{\prime *}$  from appendix A. Thus, the kinetic energy can be decomposed into two terms: a linear term  $\frac{1}{2} v^T M v$ , and an angular term  $\frac{1}{2} \omega^T I \omega$ .

#### C.2 Angular Acceleration

It is often necessary to compute  $\dot{\omega}(t)$ . Since  $L(t) = I(t)\omega(t)$ , we know  $\omega(t) = I^{-1}(t)L(t)$ . Thus,

$$\dot{\omega}(t) = \dot{I}^{-1}(t)L(t) + I^{-1}(t)\dot{L}(t).$$
 (C-4)

Since we know that  $\dot{L}(t) = \tau(t)$ , let us consider  $\dot{I}^{-1}(t)$ . From equation (2–40),

$$I^{-1}(t) = R(t)I_{body}^{-1}R(t)^{T},$$

so

$$\dot{I}^{-1}(t) = \dot{R}(t)I_{body}^{-1}R(t)^{T} + R(t)I_{body}^{-1}\dot{R}(t)^{T}.$$
 (C-5)

Since  $\dot{R}(t) = \omega(t)^* R(t)$ ,

$$\dot{R}(t)^T = (\omega(t)^* R(t))^T = R(t)^T (\omega(t)^*)^T.$$
 (C-6)

Since  $\omega(t)^*$  is antisymmetric, (i.e.  $(\omega(t)^*)^T = -\omega(t)^*$ ),

$$\dot{R}(t)^{T} = -R(t)^{T} \omega(t)^{*}. \tag{C-7}$$

This yields

$$\dot{I}^{-1}(t) = \dot{R}(t)I_{body}^{-1}R(t)^{T} + R(t)I_{body}^{-1}(-R(t)^{T}\omega(t)^{*})$$

$$= \omega(t)^{*}R(t)I_{body}^{-1}R(t)^{T} - I^{-1}(t)\omega(t)^{*}$$

$$= \omega(t)^{*}I^{-1}(t) - I^{-1}(t)\omega(t)^{*}.$$
(C-8)

Then

$$\dot{\omega}(t) = \dot{I}^{-1}(t)L(t) + I^{-1}(t)\dot{L}(t)$$

$$= \left(\omega(t)^*I^{-1}(t) - I^{-1}(t)\omega(t)^*\right)L(t) + I^{-1}(t)\dot{L}(t)$$

$$= \omega(t)^*I^{-1}(t)L(t) - I^{-1}(t)\omega(t)^*L(t) + I^{-1}(t)\dot{L}(t).$$
(C-9)

But since  $I^{-1}(t)L(t) = \omega(t)$ , the first term,  $\omega(t)^*I^{-1}(t)L(t)$  is equivalent to  $\omega(t)^*\omega(t)$ , or  $\omega(t) \times \omega(t)$ , which is zero. This leaves the final result of

$$\dot{\omega}(t) = -I^{-1}(t)\omega(t)^* L(t) + I^{-1}(t)\dot{L}(t)$$

$$= -I^{-1}(t)\omega(t) \times L(t) + I^{-1}(t)\dot{L}(t)$$

$$= I^{-1}(t)(L(t) \times \omega(t)) + I^{-1}(t)\dot{L}(t)$$

$$= I^{-1}(t)(L(t) \times \omega(t) + \dot{L}(t)).$$
(C-10)

We can see from this that even if no forces act, so that  $\dot{L}(t)$  is zero,  $\dot{\omega}(t)$  can still be non-zero. (In fact, this will happen whenever the angular momentum and angular velocities point in different directions, which in turn occurs when the body has a rotational velocity axis that is not an axis of symmetry for the body.)

#### C.3 Acceleration of a Point

Given a point of a rigid body with world space coordinate p(t), it is often necessary to compute  $\ddot{p}(t)$ . Let the body space coordinate that transforms at time t to p(t) be  $p_0$ ; then

$$p(t) = R(t) p_0 + x(t)$$

If we let r(t) = p(t) - x(t), then

$$\dot{p}(t) = \dot{R}(t)p_0 + \dot{x}(t) = \omega(t)^*R(t)p_0 + v(t)$$

$$= \omega(t) \times (R(t)p_0 + x(t) - x(t)) + v(t)$$

$$= \omega(t) \times (p(t) - x(t)) + v(t)$$

$$= \omega(t) \times r(t) + v(t).$$
(C-11)

Then

$$\ddot{p}(t) = \dot{\omega}(t) \times r(t) + \omega(t) \times \dot{r}(t) + \dot{v}(t)$$

$$= \dot{\omega}(t) \times r(t) + \omega(t) \times (\omega(t) \times r(t)) + \dot{v}(t).$$
(C-12)

We can interpret this as follows. The first term,  $\dot{\omega}(t) \times r(t)$  is the *tangential* acceleration of the point; that is,  $\dot{\omega}(t) \times r(t)$  is the acceleration perpendicular to the displacement r(t) as a result of the body being angularly accelerated. The second term,  $\omega(t) \times (\omega(t) \times r(t))$  is the centripetal acceleration of the point; this centripetal acceleration arises because the body is rigid, and points on the body must rotate in a circular orbit about the center of mass. The last term,  $\dot{v}(t)$  is the linear acceleration of the point due to the linear acceleration of the center of mass of the body.

# **Appendix D** Resting Contact Derivations

If you're determined to implement resting contact in your simulator, you'll need the derivations and the code in this appendix. This is probably not a fun appendix to work through; then again, this wasn't a fun appendix to write! The derivations in here are somewhat terse, but the code at the end of the appendix will hopefully make things clearer.

#### **D.1** Derivations

We need to express  $\ddot{d}_i(t_0)$  in terms of all the unknown  $f_i$ 's. It will turn out that we'll be able to write each  $\ddot{d}_i(t_0)$  in the form

$$\ddot{d}_i(t_0) = a_{i1}f_1 + a_{i2}f_2 + \dots + a_{in}f_n + b_i.$$
 (D-1)

Given i and j, we need to know how  $\ddot{d}_i(t_0)$  depends on  $f_j$ , that is, we need to know  $a_{ij}$ . Also, we need to compute the constant term  $b_i$ .

Let's start by determining  $a_{ij}$  and ignoring the constant part  $b_i$ . We'll assume the *i*th contact involves two bodies A and B. From equation (9–4), we can write  $d_i(t_0)$  as

$$\ddot{d}(t_0) = \hat{n}_i(t_0) \cdot (\ddot{p}_a(t_0) - \ddot{p}_b(t_0)) + 2\dot{\hat{n}}_i(t_0) \cdot (\dot{p}_a(t_0) - \dot{p}_b(t_0)) \tag{D-2}$$

where  $p_a(t_0) = p_i = p_b(t_0)$  is the contact point for the *i*th contact at time  $t_0$ . The right-most term  $2\hat{n}_i(t_0) \cdot (\dot{p}_a(t_0) - \dot{p}_b(t_0))$  is a velocity dependent term (i.e. you can immediately calculate it without knowing the forces involved), and is part of  $b_i$ , so we'll ignore this for now.

So we only need to know how  $\ddot{p}_a(t_0)$  and  $\ddot{p}_b(t_0)$  depend on  $f_j$ , the magnitude of the jth contact force. Consider the jth contact. If body A is not one of the bodies involved in the jth contact, then  $\ddot{p}_a(t_0)$  is independent of  $f_j$ , because the jth contact force does not act on body A. Similarly, if B is also not one of the two bodies involved in the jth contact, then  $\ddot{p}_b(t_0)$  is also independent of  $f_j$ . (For example, in figure 26, the acceleration of the contact points at the first contact is completely unaffected by the contact force acting at the fifth contact. Thus,  $\ddot{d}_1(t_0)$  would be completely independent of  $f_5$ . Conversely,  $\ddot{d}_5(t_0)$  is completely independent of  $f_1$ .)

Suppose though that in the *j*th contact, body *A* is involved. For definiteness, suppose that in the *j*th contact, a force of  $j\hat{n}_j(t_0)$  acts on body *A*, as opposed to  $-j\hat{n}_j(t_0)$ . Let's derive how  $\ddot{p}_a(t_0)$  is affected by the force  $j\hat{n}_j(t_0)$  acting on *A*.

From equation (C-12), we can write

$$\ddot{p}_a(t) = \dot{v}_a(t) + \dot{\omega}_a(t) \times r_a(t) + \omega_a(t) \times (\omega_a(t) \times r_a(t)) \tag{D-3}$$

where  $r_a(t) = p_a(t) - x_a(t)$ , and  $x_a(t)$ ,  $v_a(t)$ , and  $\omega_a(t)$  are all the variables associated with body A. We know that  $\dot{v}_a(t)$  is the linear acceleration of body A, and is equal to the total force acting on A divided by the mass. Thus, a force of  $j\hat{n}_i(t_0)$  contributes

$$\frac{f_j \hat{n}_j(t_0)}{m_a} = f_j \frac{\hat{n}_j(t_0)}{m_a}$$
 (D-4)

to  $\dot{v}_a(t)$  and thus  $\ddot{p}_a(t)$ . Similarly, consider  $\dot{\omega}_a(t)$ , from equation (C–10):

$$\dot{\omega}_a(t) = I_a^{-1}(t)\tau_a(t) + I_a^{-1}(t)(L_a(t) \times \omega_a(t))$$

where  $\tau_a(t)$  is the total torque acting on body A. If the jth contact occurs at the point  $p_j$ , then the force  $j\hat{n}_j(t_0)$  exerts a torque of

$$(p_i - x_a(t_0)) \times f_i \hat{n}_i(t_0).$$

Thus, the angular contribution to  $\ddot{p}_a(t_0)$  is

$$f_j\left(I_a^{-1}(t_0)\left((p_j-x_a(t_0))\times\hat{n}_j(t_0)\right)\right)\times r_a. \tag{D-5}$$

The total dependence of  $\ddot{p}_a(t_0)$  on  $f_i$  is therefore

$$f_j\left(\frac{\hat{n}_j(t_0)}{m_a} + \left(I_a^{-1}(t_0)\left((p_j - x_a(t_0)) \times \hat{n}_j(t_0)\right)\right) \times r_a\right).$$

Now, if a force of  $-f_j\hat{n}(t_0)$  had acted on A instead, we'd get the same dependence, but with a minus sign in front of  $f_j$ . Clearly,  $\ddot{p}_b(t_0)$  depends on  $f_j$  in the same sort of manner. Once we compute how  $\ddot{p}_a(t_0)$  and  $\ddot{p}_b(t_0)$  depend on  $f_j$ , we combine the results together and take the dot product with  $\hat{n}_i(t_0)$ , to see how  $\ddot{d}_i(t_0)$  depends on  $f_j$ . This gives us  $a_{ij}$ . Confused? See the code below.

We still need to compute  $b_i$ . We know that  $d_i(t_0)$  contains the constant term

$$2\dot{\hat{n}}_i(t_0) \cdot (\dot{p}_a(t_0) - \dot{p}_b(t_0)).$$

But we also have to take into account the contributions to  $\ddot{p}_a(t_0)$  and  $\ddot{p}_b(t_0)$  due to known external forces such as gravity, as well as the force-independent terms  $\omega_a(t_0) \times (\omega_a(t_0) \times r_a)$  and  $(I_a^{-1}(t_0)(\pounds_a(t_0) \times \omega_a(t_0))) \times r_a$ . If we let the net external force acting on A be  $F_a(t_0)$ , and the net external torque be  $\tau_a(t_0)$ , then from equations (D-4) and (D-5), we get that  $F_a(t_0)$  contributes

$$\frac{F_a(t_0)}{m_a}$$

and that  $\tau_a(t_0)$  contributes

$$\left(I_a^{-1}(t_0)\tau_a(t_0)\right)\times r_a.$$

Thus, the part of  $\ddot{p}_a(t_0)$  that is independent from all the  $f_i$ 's is

$$\frac{F_a(t_0)}{m_a} + \left(I_a^{-1}(t_0)\tau_a(t_0)\right) \times r_a + \omega_a(t_0) \times (\omega_a(t_0) \times r_a) + \left(I_a^{-1}(t_0)(\mathbb{E}_a(t_0) \times \omega_a(t_0))\right) \times r_a$$

and similarly for  $\ddot{p}_b(t_0)$ . To compute  $b_i$ , we combine the constant parts of  $\ddot{p}_a(t_0)$ ,  $\ddot{p}_b(t_0)$ , dot with  $\hat{n}_i(t_0)$ , and add the term  $2\dot{\hat{n}}_i(t_0) \cdot (\dot{p}_a(t_0) - \dot{p}_b(t_0))$ .

#### D.2 Code

Here's the code to implement the above derivations. Let's start by computing the constant  $b_i$  terms.

```
/* return the derivative of the normal vector */
triple computeNdot(Contact *c)
    if(c->vf)
                  /* vertex/face contact */
        /* The vector 'n' is attached to B, so... */
        return c->b->omega ^ c->n;
    else
    {
         /* This is a little trickier. The unit normal `n' is
                \hat{n} = \frac{\text{ea} \times \text{eb}}{\|\text{ea} \times \text{eb}\|}
            Differentiating \hat{n} with respect to time is left
            as an exercise... but here's some code */
        triple eadot = c->a->omega ^ ea,
                                                  /* ė<sub>a</sub> */
                                                 /* ė<sub>h</sub> */
                 ebdot = c->b->omega ^ eb;
                 n1 = ea * eb,
                 z = eadot * eb + ea * ebdot;
        double l = length(n1);
                                        /* normalize */
        n1 = n1 / length;
        return (z - ((z * n) * n)) / l;
    }
}
void compute_b_vector(Contact contacts[], int ncontacts, vector &b)
{
    for(int i = 0; i < ncontacts; i++)</pre>
        Contact *c = &contacts[i];
        Body
                 *A = c->a
                 *B = c->b;
        triple n = c->n, /* \hat{n}_i(t_0) */
                 ra = c->p - A->x,
                                       /* p - x_a(t_0) * /
                 rb = c->p - B->x; /* p-x_h(t_0) */
         /* Get the external forces and torques */
        triple f_ext_a = A->force,
                 f_ext_b = B->force,
                 t_ext_a = A->torque,
```

```
t_ext_b = B->torque;
        triple a_ext_part, a_vel_part,
                b_ext_part, b_vel_part;
        /* Operators: `^' is for cross product, `*', is for
           dot products (between two triples), or matrix-vector
           multiplication (between a matrix and a triple). */
        /* Compute the part of \ddot{p}_a(t_0) due to the external
           force and torque, and similarly for \ddot{p}_h(t_0). */
        a_ext_part = f_ext_a / A->mass + ((A->Iinv * t_ext_a) ^ ra),
        b_ext_part = f_ext_b / B->mass + ((B->linv * t_ext_b) ^ rb);
        /* Compute the part of \ddot{p}_a(t_0) due to velocity,
           and similarly for \ddot{p}_b(t_0). */
        a_vel_part = (A->omega ^ (A->omega ^ ra)) +
                 ((A->Iinv * (A->L ^ A->omega)) ^ ra);
        b_vel_part = (B->omega ^ (B->omega ^ rb)) +
                 ((B->Iinv * (B->L ^ B->omega)) ^ rb);
        /* Combine the above results, and dot with \hat{n}_i(t_0) */
        double k1 = n * ((a_ext_part + a_vel_part) -
                           (b_ext_part + b_vel_part));
        triple ndot = computeNdot(c);
        /* See section 8 for 'pt_velocity' definition */
        double k2 = 2 * ndot * (pt_velocity(A, c->p) -
                                  pt_velocity(B, c->p));
        b[i] = k1 + k2;
    }
}
```

Computing the  $a_{ij}$  terms is a little more tricky, because we have to keep track of how the jth contact force affects the ith contact point. The following routine is not the most efficient way to do things, because with a good data structure, you can tell in advance which of the  $a_{ij}$ 's are going to be zero. Still unless you're working with really huge numbers of contacts, not too much extra work will be done.

```
void compute_a_matrix(Contact contacts[], int ncontacts, bigmatrix &a)
    for(int i = 0; i < ncontacts; i++)</pre>
        for(int j = 0; j < ncontacts; j++)</pre>
            a[i,j] = compute aij(contacts[i], contacts[j]);
}
double compute_aij(Contact ci, Contact cj)
    /* If the bodies involved in the ith and jth contact are
       distinct, then a_{ij} is zero. */
    if((ci.a != cj.a) && (ci.b != cj.b) &&
       (ci.a != cj.b) && (ci.b != cj.a))
        return 0.0;
           *A = ci.a,
    Body
           *B = ci.b;
                             /* \hat{n}_i(t_0) */
    triple ni = ci.n,
                             /* \hat{n}_{i}(t_{0}) * /
           nj = cj.n,
           pi = ci.p,
                             /* ith contact point location */
                              /* jth contact point location */
           pj = cj.p,
           ra = pi - A->x,
           rb = pi - B -> x;
    /* What force and torque does contact j exert on body A? */
    triple force_on_a = 0,
           torque_on_a = 0;
    if(cj.a == ci.a)
        /* force direction of jth contact force on A */
        force_on_a = nj;
        /* torque direction */
        torque_on_a = (pj - A->x) ^ nj;
    }
    else if(cj.b == ci.a)
        force_on_a = - nj;
        torque_on_a = (pj - A->x) ^ nj;
    }
```

```
/* What force and torque does contact i exert on body B? */
triple force_on_b = 0,
       torque_on_b = 0;
if(cj.a == ci.b)
    /* force direction of jth contact force on B */
    force_on_b = nj;
    /* torque direction */
    torque_on_b = (pj - B->x) ^ nj;
else if(cj.b == ci.b)
    force_on_b = - nj;
    torque\_on\_b = (pj - B->x) ^ nj;
}
/st Now compute how the jth contact force affects the linear
   and angular acceleration of the contact point on body A */
triple a_linear = force_on_a / A->mass,
       a_angular = (A->Iinv * torque_on_a) ^ ra;
/* Same for B */
triple b_linear = force_on_b / B->mass,
       b_angular = (B->Iinv * torque_on_b) ^ rb;
return ni * ((a_linear + a_angular) - (b_linear + b_angular));
```

}

### **References**

- [1] D. Baraff. Analytical methods for dynamic simulation of non-penetrating rigid bodies. In *Computer Graphics (Proc. SIGGRAPH)*, volume 23, pages 223–232. ACM, July 1989.
- [2] D. Baraff. Curved surfaces and coherence for non-penetrating rigid body simulation. In *Computer Graphics (Proc. SIGGRAPH)*, volume 24, pages 19–28. ACM, August 1990.
- [3] D. Baraff. Fast contact force computation for nonpenetrating rigid bodies. *Computer Graphics* (*Proc. SIGGRAPH*), 28:23–34, 1994.
- [4] J. Canny. Collision detection for moving polyhedra. *IEEE Transactions on Pattern Analysis and Machine Intelligence*, 8(2), 1986.
- [5] P.A. Cundall. Formulation of a three-dimensional distinct element model—Part I. A scheme to represent contacts in a system composed of many polyhedral blocks. *International Journal of Rock Mechanics, Mineral Science and Geomechanics*, 25, 1988.
- [6] E.G. Gilbert and S.M. Hong. A new algorithm for detecting the collision of moving objects. In *International Conference on Robotics and Automation*, pages 8–13. IEEE, 1989.
- [7] P. Gill, S. Hammarling, W. Murray, M. Saunders, and M. Wright. User's guide for LSSOL: A Fortran package for constrained linear least-squares and convex quadratic programming. Technical Report Sol 86-1, Systems Optimization Laboratory, Department of Operations Research, Stanford University, 1986.
- [8] P. Gill, W. Murray, M. Saunders, and M. Wright. User's guide for QPSOL: A Fortran package for quadratic programming. Technical Report Sol 84-6, Systems Optimization Laboratory, Department of Operations Research, Stanford University, 1984.
- [9] P. Gill, W. Murray, M. Saunders, and M. Wright. User's guide for NPSOL: A Fortran package for nonlinear programming. Technical Report Sol 86-2, Systems Optimization Laboratory, Department of Operations Research, Stanford University, 1986.
- [10] H. Goldstein. Classical Mechanics. Addison-Wesley, Reading, 1983.
- [11] W. Meyer. Distance between boxes: Applications to collision detection and clipping. In *International Conference on Robotics and Automation*, pages 597–602. IEEE, 1986.
- [12] P.M. Moore and J. Wilhelms. Collision detection and reponse for computer animation. In *Computer Graphics (Proc. SIGGRAPH)*, volume 22, pages 289–298. ACM, August 1988.
- [13] F.P. Preparata and M.I. Shamos. Computational Geometry. Springer-Verlag, New York, 1985.
- [14] W.H. Press, B.P. Flannery, S.A. Teukolsky, and W.T. Vetterling. *Numerical Recipes*. Cambridge University Press, 1986.
- [15] R. Sedgewick. Algorithms. Addison-Wesley, 1983.
- [16] K. Shoemake. Animating rotation with quaternion curves. In *Computer Graphics (Proc. SIGGRAPH)*, volume 19, pages 245–254. ACM, July 1985.
- [17] B. Von Herzen, A. Barr, and H. Zatz. Geometric collisions for time-dependent parametric surfaces. In *Computer Graphics (Proc. SIGGRAPH)*, volume 24, pages 39–48. ACM, August 1990.