# Specifications of FDPS

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# Contents

| 1        | Cor  | ncept of Document   | 15 |
|----------|------|---|----|
| <b>2</b> | Cor  | ncept of FDPS   | 16 |
|          | 2.1  | Purpose of the development of FDPS                          | 16 |
|          | 2.2  | Basic concept   | 16 |
|          |      | 2.2.1 Procedures of massively parallel particle simulations | 16 |
|          |      | 2.2.2 Division of tasks between users and FDPS              | 17 |
|          |      | 2.2.3 Users' tasks  | 17 |
|          |      | 2.2.4 Complement  | 17 |
|          | 2.3  | The structure of a simulation code with FDPS                | 18 |
|          | ۷.5  | The structure of a simulation code with PDI 5               | 10 |
| 3        | Fil€ | es and directories  | 19 |
|          | 3.1  | Summary   | 19 |
|          | 3.2  | Documents   | 19 |
|          | 3.3  | Source files  | 19 |
|          | 0.0  | 3.3.1 Extended features                                     | 19 |
|          |      | 3.3.1.1 Particle Mesh scheme                                | 20 |
|          |      | 3.3.1.2 x86 version Phantom-GRAPE                           | 20 |
|          |      | 3.3.1.2.1 Low precision                                     | 20 |
|          |      | <del>-</del>  |    |
|          |      | 3.3.1.2.2 Low precision with cutoff                         | 20 |
|          |      | 3.3.1.2.3 High precision                                    | 20 |
|          | 3.4  | Test codes  | 20 |
|          | 3.5  | Sample codes  | 20 |
|          |      | 3.5.1 Gravitational $N$ -body simulation                    | 20 |
|          |      | 3.5.2 SPH Simulation  | 21 |

| 4 | Mae | cros at | compile 1   | $_{ m time}$ |          |               |     |  |      |   |   |  |  |  | 22 |
|---|-----|---------|-------------|--------------|----------|---------------|-----|--|------|---|---|--|--|--|----|
|   | 4.1 | Summa   | ary         |              |          |               |     |  | <br> |   |   |  |  |  | 22 |
|   | 4.2 | Coordi  | nate system | m            |          |               |     |  | <br> |   |   |  |  |  | 22 |
|   |     | 4.2.1   | Summary     |              |          |               |     |  | <br> |   |   |  |  |  | 22 |
|   |     | 4.2.2   | 3D Cartes   | sian coord   | dinate s | ystem         | ١   |  | <br> |   |   |  |  |  | 22 |
|   |     | 4.2.3   | 2D Cartes   | sian coord   | dinate s | $_{ m ystem}$ | ١   |  | <br> |   |   |  |  |  | 22 |
|   | 4.3 | Paralle | l processin |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 4.3.1   | Summary     | ~            |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 4.3.2   | OpenMP      |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 4.3.3   | MPI         |              |          |               |     |  | <br> |   |   |  |  |  | 22 |
|   | 4.4 | Accura  | cy of data  | types.       |          |               |     |  | <br> |   |   |  |  |  | 22 |
|   |     | 4.4.1   | Summary     |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 4.4.2   | Accuracy    |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     |         | v           | 1            |          |               |     |  |      | • | • |  |  |  |    |
| 5 | Nar | nespac  |             |              |          |               |     |  |      |   |   |  |  |  | 23 |
|   | 5.1 |         | ary         |              |          |               |     |  |      |   |   |  |  |  |    |
|   | 5.2 | Particl | eSimulator  | ·            |          |               |     |  | <br> |   |   |  |  |  | 23 |
|   |     | 5.2.1   | ParticleM   | esh          |          |               |     |  | <br> |   |   |  |  |  | 23 |
| _ |     | _       |             |              |          |               |     |  |      |   |   |  |  |  |    |
| 6 |     | a Type  |             |              |          |               |     |  |      |   |   |  |  |  | 24 |
|   | 6.1 |         | ary         |              |          |               |     |  |      |   |   |  |  |  |    |
|   | 6.2 | _       | type        |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 6.2.1   | Overview    |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 6.2.2   | PS::S32     |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 6.2.3   | PS::S64     |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 6.2.4   | PS::U32     |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     | 6.2.5   | PS::U64     |              |          |               |     |  |      |   |   |  |  |  |    |
|   | 6.3 | Floatin | ig point ty | ре           |          |               |     |  | <br> |   |   |  |  |  | 25 |
|   |     | 6.3.1   | Abstract    |              |          |               |     |  | <br> |   |   |  |  |  | 25 |
|   |     | 6.3.2   | PS::F32     |              |          |               |     |  | <br> |   |   |  |  |  | 25 |
|   |     | 6.3.3   | PS::F64     |              |          |               |     |  | <br> |   |   |  |  |  | 25 |
|   | 6.4 | Vector  | type        |              |          |               |     |  | <br> |   |   |  |  |  | 25 |
|   |     | 6.4.1   | Abstract    |              |          |               |     |  | <br> |   |   |  |  |  | 25 |
|   |     | 6.4.2   | PS::Vecto   | r2           |          |               |     |  | <br> |   |   |  |  |  | 25 |
|   |     |         | 6.4.2.1     | Construct    | tor      |               |     |  | <br> |   |   |  |  |  | 27 |
|   |     |         | 6.4.2.2     | Copy con     | structor |               |     |  | <br> |   |   |  |  |  | 27 |
|   |     |         | 6.4.2.3     | Member v     | variable | s             |     |  | <br> |   |   |  |  |  | 28 |
|   |     |         | 6.4.2.4     | Assignme     | nt oper  | ator .        |     |  | <br> |   |   |  |  |  | 28 |
|   |     |         | 6.4.2.5     | operato      | rs       |               |     |  | <br> |   |   |  |  |  | 28 |
|   |     |         | 6.4.2.6     | Addition     | and sub  | otract        | ion |  | <br> |   |   |  |  |  | 29 |
|   |     |         |             | Scalar mu    |          |               |     |  |      |   |   |  |  |  |    |
|   |     |         |             | Inner and    | -        |               |     |  |      |   |   |  |  |  |    |
|   |     |         |             | Typecast     | -        |               |     |  |      |   |   |  |  |  |    |
|   |     | 6.4.3   | PS::Vecto   |              |          |               |     |  |      |   |   |  |  |  |    |
|   |     |         |             | Construct    |          |               |     |  |      |   |   |  |  |  |    |
|   |     |         |             | Copy con     |          |               |     |  |      |   |   |  |  |  |    |

|     |               | 6.4.3.3         | Member variables               |
|-----|---------------|-----------------|--------------------------------|
|     |               | 6.4.3.4         | Assignment operator            |
|     |               | 6.4.3.5         | [] operators                   |
|     |               | 6.4.3.6         | Addition and subtraction       |
|     |               | 6.4.3.7         | Scalar multiplication          |
|     |               | 6.4.3.8         | Inner and outer product        |
|     |               | 6.4.3.9         | Typecast to Vector3 <u></u>    |
|     | 6.4.4         | Wrappe          | rs                             |
| 6.5 | Ortho         | tope type       | 9                              |
|     | 6.5.1         | Abstrac         | t                              |
|     | 6.5.2         | PS::Ort         | hotope2                        |
|     |               | 6.5.2.1         | Member variables               |
|     |               | 6.5.2.2         | Constructors                   |
|     |               | 6.5.2.3         | Copy constructor               |
|     |               | 6.5.2.4         | Initialize                     |
|     |               | 6.5.2.5         | Merge operations               |
|     | 6.5.3         | PS::Ort         | hotope3                        |
|     |               | 6.5.3.1         | Member variables               |
|     |               | 6.5.3.2         | Constructors                   |
|     |               | 6.5.3.3         | Copy constructor               |
|     |               | 6.5.3.4         | Initialize                     |
|     |               | 6.5.3.5         | Merge operations               |
|     | 6.5.4         | Wrappe          | 0 1                            |
| 6.6 |               |                 | rix type                       |
| 0.0 | 6.6.1         |                 | t                              |
|     | 6.6.2         |                 | rixSym2                        |
|     | 0.0.2         | 6.6.2.1         | Constructor                    |
|     |               | 6.6.2.2         | Copy constructor               |
|     |               | 6.6.2.3         | Assignment operator            |
|     |               | 6.6.2.4         | Addition and subtraction       |
|     |               | 6.6.2.5         | Trace                          |
|     |               | 6.6.2.6         | Typecast to MatrixSym2 <u></u> |
|     | 6.6.3         |                 | rixSym3                        |
|     | 0.0.5         | 6.6.3.1         | Constructor                    |
|     |               | 6.6.3.2         | Copy constructor               |
|     |               | 6.6.3.3         | Assignment operator            |
|     |               | 6.6.3.4         | Addition and subtraction       |
|     |               | 6.6.3.5         | Trace                          |
|     |               | 6.6.3.6         | Typecast to MatrixSym3 <u></u> |
|     | 6.6.4         | Wrappe          | v -                            |
| 6.7 |               |                 | rs                             |
| 0.7 | 6.7.1         |                 | V 1                            |
|     | 6.7.1         | Summai<br>PSSEA | ry                             |
|     | 6.7.2 $6.7.3$ |                 | ARCH_MODE_LONG                 |
|     | 6.7.3 $6.7.4$ |                 | ARCH_MODE_GATHER               |
|     | 6.7.4 $6.7.5$ |                 |                                |
|     | 0.1.0         | F O O D P       | ARCH_MODE_SCATTER              |

|   |     | 6.7.6        |   | 59  |
|---|-----|--------------|---|-----|
|   |     | 6.7.7        | PS::SEARCH_MODE_LONG_SCATTER                                      | 59  |
|   |     | 6.7.8        | PS::SEARCH_MODE_LONG_SYMMETRY                                     | 59  |
|   |     | 6.7.9        | PS::SEARCH_MODE_LONG_CUTOFF_SCATTER                               | 59  |
|   | 6.8 | Enum         | erated type   | 59  |
|   |     | 6.8.1        | Summary   | 59  |
|   |     | 6.8.2        | PS::BOUNDARY_CONDITION type                                       | 59  |
|   |     |              |   | 59  |
|   |     |              | 6.8.2.2 PS::BOUNDARY_CONDITION_OPEN                               | 60  |
|   |     |              | 6.8.2.3 PS::BOUNDARY_CONDITION_PERIODIC_X                         | 60  |
|   |     |              | 6.8.2.4 PS::BOUNDARY_CONDITION_PERIODIC_Y                         | 60  |
|   |     |              | 6.8.2.5 PS::BOUNDARY_CONDITION_PERIODIC_Z                         | 60  |
|   |     |              | 6.8.2.6 PS::BOUNDARY_CONDITION_PERIODIC_XY                        | 60  |
|   |     |              | 6.8.2.7 PS::BOUNDARY_CONDITION_PERIODIC_XZ                        | 60  |
|   |     |              | 6.8.2.8 PS::BOUNDARY_CONDITION_PERIODIC_YZ                        | 61  |
|   |     |              | 6.8.2.9 PS::BOUNDARY_CONDITION_PERIODIC_XYZ                       | 61  |
|   |     |              | 6.8.2.10 PS::BOUNDARY_CONDITION_SHEARING_BOX                      | 61  |
|   |     |              | 6.8.2.11 PS::BOUNDARY_CONDITION_USER_DEFINED                      | 61  |
|   |     | 6.8.3        | PS::INTERACTION_LIST_MODE type                                    | 61  |
|   |     |              |   | 61  |
|   |     |              | 6.8.3.2 PS::MAKE_LIST   | 61  |
|   |     |              | 6.8.3.3 PS::MAKE_LIST_FOR_REUSE                                   | 61  |
|   |     |              | 6.8.3.4 PS::REUSE_INTERACTION_LIST                                | 62  |
|   | 6.9 | PS::T        | imeProfile type   | 62  |
|   |     | 6.9.1        | Abstract  | 62  |
|   |     |              | 6.9.1.1 Addition  | 62  |
|   |     |              | 6.9.1.2 Reduction   | 63  |
|   |     |              | 6.9.1.3 Initialize  | 63  |
| _ | ~1  |              |   |     |
| 7 |     |              |   | 64  |
|   | 7.1 |              | v   | 64  |
|   | 7.2 |              | article   |     |
|   |     | 7.2.1        | V   | 64  |
|   |     | 7.2.2        |   | 64  |
|   |     | 7.2.3        | 1   | 65  |
|   |     |              | v   | 65  |
|   |     |              | 8   | 65  |
|   |     | <b>-</b> 2 4 | 10  | 65  |
|   |     | 7.2.4        | 1   | 66  |
|   |     |              | v   | 66  |
|   |     |              | 7.2.4.2 Modes other than PS::SEARCH_MODE_LONG for PS::SEARCH_MODE | 0.0 |
|   |     |              |   | 66  |
|   |     |              | O   | 66  |
|   |     |              | , · · · · · · · · · · · · · · · · · · ·                           | 66  |
|   |     |              |   | 67  |
|   |     |              | 7.2.4.3.2 FP::writeAscii  | 67  |

|     |       | 7.2.4.3.3 FP::readBinary 6   |
|-----|-------|--|
|     |       | 7.2.4.3.4 FP::writeBinary 6  |
|     |       | 7.2.4.4 ParticleSystem::adjustPositionIntoRootDomain API is  |
|     |       | $used \dots \dots$ |
|     |       | 7.2.4.4.1 FP::setPos 6   |
|     |       | 7.2.4.5 Particle Mesh class, which is an extension of FDPS, is used 6  |
|     |       | 7.2.4.5.1 FP::getChargeParticleMesh 6  |
|     |       | 7.2.4.5.2 FP::copyFromForceParticleMesh 6  |
| 7.3 | Essen | tialParticleI  |
|     | 7.3.1 | Summary  |
|     | 7.3.2 | Premise  |
|     | 7.3.3 | Required member functions  |
|     |       | 7.3.3.1 Summary  |
|     |       | 7.3.3.2 EPI::getPos  |
|     |       | 7.3.3.3 EPI::copyFromFP  |
|     | 7.3.4 | Required member functions for specific case  |
|     |       | 7.3.4.1 Summary  |
|     |       | 7.3.4.2 Modes other than PS::SEARCH_MODE_LONG as PS::SEARCH_MODE   |
|     |       | are used   |
|     |       | 7.3.4.2.1 EPI::getRSearch  |
| 7.4 | Essen | tialParticleJ  |
|     | 7.4.1 | Summary  |
|     | 7.4.2 | Premise  |
|     | 7.4.3 | Required member functions  |
|     |       | 7.4.3.1 Summary  |
|     |       | 7.4.3.2 EPJ::getPos  |
|     |       | 7.4.3.3 EPJ::copyFromFP  |
|     | 7.4.4 | Required member functions for specific cases   |
|     |       | 7.4.4.1 Summary  |
|     |       | 7.4.4.2 Modes other than PS::SEARCH_MODE_LONG as PS::SEARCH_MODE   |
|     |       | are used   |
|     |       | 7.4.4.2.1 EPJ::getRSearch  |
|     |       | 7.4.4.3 Modes other than PS::BOUNDARY_CONDITION_OPEN are used  |
|     |       | BOUNDARY_CONDITION   |
|     |       | 7.4.4.3.1 EPJ::setPos  |
|     |       | 7.4.4.4 Obtian EPJ from id of particle   |
|     |       | 7.4.4.4.1 EPJ::getId   |
| 7.5 | Mome  | ent  |
|     | 7.5.1 | Summary  |
|     | 7.5.2 | Pre-defined class  |
|     |       | 7.5.2.1 Summary  |
|     |       | 7.5.2.2 PS::SEARCH_MODE_LONG   |
|     |       | 7.5.2.2.1 PS::MomentMonopole   |
|     |       | 7.5.2.2.2 PS::MomentQuadrupole   |
|     |       | 7.5.2.2.3 PS::MomentMonopoleGeometricCenter  |
|     |       | 7.5.2.2.4 PS::MomentDipoleGeometricCenter  |

|     |       | 7.5.2.2.5 PS::MomentQuadrupoleGeometricCenter 7 |
|-----|-------|---|
|     |       | 7.5.2.3 PS::SEARCH_MODE_LONG_SCATTER            |
|     |       | 7.5.2.3.1 PS::MomentMonopoleScatter             |
|     |       | 7.5.2.3.2 PS::MomentQuadrupoleScatter           |
|     |       | 7.5.2.4 PS::SEARCH_MODE_LONG_SYMMETRY           |
|     |       | 7.5.2.4.1 PS::MomentMonopoleSymmetry            |
|     |       | 7.5.2.4.2 PS::MomentQuadrupoleSymmetry 8        |
|     |       | 7.5.2.5 PS::SEARCH_MODE_LONG_CUTOFF 8           |
|     |       | 7.5.2.5.1 PS::MomentMonopoleCutoff 8            |
|     | 7.5.3 | Required member functions                       |
|     |       | 7.5.3.1 Summary                                 |
|     |       | 7.5.3.2 Constructor                             |
|     |       | 7.5.3.3 Mom::init                               |
|     |       | 7.5.3.4 Mom::getPos                             |
|     |       | 7.5.3.5 Mom::getCharge                          |
|     |       | 7.5.3.6 Mom::accumulateAtLeaf                   |
|     |       | 7.5.3.7 Mom::accumulate                         |
|     |       | 7.5.3.8 Mom::set                                |
|     |       | 7.5.3.9 Mom::accumulateAtLeaf2                  |
|     |       | 7.5.3.10 Mom::accumulate2                       |
|     | 7.5.4 | Required member functions for specific cases    |
|     |       | 7.5.4.1 Summary                                 |
|     |       | 7.5.4.2 One of PS::SEARCH_MODE_LONG_CUTOFF,     |
|     |       | PS::SEARCH_MODE_LONG_SCATTER,                   |
|     |       | PS::SEARCH_MODE_LONG_SYMMETRY                   |
|     |       | is used as PS::SEARCH_MODE                      |
|     |       | 7.5.4.3 Mom::getVertexIn                        |
|     |       | 7.5.4.4 Mom::getVertexOut                       |
| 7.6 | Super | ParticleJ                                       |
|     | 7.6.1 | Summary   |
|     | 7.6.2 | Pre-defined class                               |
|     |       | 7.6.2.1 PS::SEARCH_MODE_LONG                    |
|     |       | 7.6.2.1.1 PS::SPJMonopole                       |
|     |       | 7.6.2.1.2 PS::SPJQuadrupole 8                   |
|     |       | 7.6.2.1.3 PS::SPJMonopoleGeometricCenter 89     |
|     |       | 7.6.2.1.4 PS::SPJDipoleGeometricCenter 89       |
|     |       | 7.6.2.1.5 PS::SPJQuadrupoleGeometricCenter 9    |
|     |       | 7.6.2.2 PS::SEARCH_MODE_LONG_SCATTER 9          |
|     |       | 7.6.2.2.1 PS::SPJMonopoleScatter                |
|     |       | 7.6.2.2.2 PS::SPJQuadrupoleScatter 9            |
|     |       | 7.6.2.3 PS::SEARCH_MODE_LONG_SYMMETRY 9         |
|     |       | 7.6.2.3.1 PS::SPJMonopoleSymmetry 9             |
|     |       | 7.6.2.3.2 PS::SPJQuadrupoleSymmetry 9           |
|     |       | 7.6.2.4 PS::SEARCH_MODE_LONG_CUTOFF 9           |
|     |       | 7.6.2.4.1 PS::SPJMonopoleCutoff                 |
|     | 7.6.3 | Required member functions                       |

|      |               | 7.6.3.1    | Summary                              | 93          |
|------|---------------|------------|--------------------------------------|-------------|
|      |               | 7.6.3.2    | SPJ::getPos                          | 93          |
|      |               | 7.6.3.3    | SPJ::setPos                          | 94          |
|      |               | 7.6.3.4    | SPJ::copyFromMoment                  | 94          |
|      |               | 7.6.3.5    | SPJ::convertToMoment                 | 95          |
|      |               | 7.6.3.6    | SPJ::clear                           | 95          |
| 7.7  | Force         |            |                                      | 95          |
|      | 7.7.1         |            | у                                    | 95          |
|      | 7.7.2         |            |                                      | 96          |
|      | 7.7.3         |            | d member functions                   | 96          |
|      | 1.1.0         |            | Result::clear                        | 96          |
| 7.8  | Heade         |            |                                      | 96          |
| 1.0  | 7.8.1         |            | у                                    | 96          |
|      | 7.8.2         |            |                                      | 96          |
|      | 7.8.3         |            | l member functions for specific case | 97          |
|      | 1.0.0         | 7.8.3.1    | Hdr::readAscii                       | 97          |
|      |               | 7.8.3.2    | Hdr::writeAscii                      | 97          |
|      |               | 7.8.3.3    | Hdr::readBinary                      | 97          |
|      |               | 7.8.3.4    | Hdr::writeBinary                     | 98          |
| 7.9  | Functo        |            | ceEpEp                               | 98          |
| 1.9  | 7.9.1         |            | у                                    | 98          |
|      | 7.9.1 $7.9.2$ |            | y                                    | 98          |
|      | 7.9.2 $7.9.3$ |            | pEp::operator ()                     | 98          |
| 7 10 |               |            | се $\operatorname{SpEp}$             | 90          |
| 1.10 |               |            | у                                    | 99          |
|      |               |            | y                                    | 99          |
|      |               |            | pEp::operator ()                     | 99          |
| 7 11 |               |            | ceDispatch $\dots$                   | 100         |
| 1.11 |               |            | у                                    | 100         |
|      |               |            | e of short-range interactions        | 100         |
|      |               |            | long-range interaction               | 101         |
| 7 19 |               |            | ceRetrieve                           |             |
| 1.12 |               |            | у                                    |             |
|      | 1.12.1        | Summar     | y                                    | 102         |
| Init | ializati      | on and     | Finalization of FDPS                 | <b>10</b> 4 |
| 8.1  |               |            |                                      | 104         |
| 8.2  |               |            |                                      | 104         |
|      | 8.2.1         |            |                                      | 104         |
|      | 8.2.2         | PS::Fina   | dize                                 | 104         |
|      | 8.2.3         |            | rt                                   | 105         |
|      | 8.2.4         |            |                                      | 105         |
|      |               |            |                                      |             |
| Clas |               |            |                                      | 106         |
| 9.1  | Standa        | ard Classe | es                                   | 106         |
|      | 9.1.1         | Summar     | у                                    | 106         |
|      | 9.1.2         |            |                                      | 106         |

|       | 9.1.2.1 Creation | n of Object  | 106    |
|-------|------------------|--|--------|
|       |                  |  | 106    |
|       | 9.1.2.2.1        | Initial Setup  | 106    |
|       | 9.1.2.2.1.1      | Constructor  | 107    |
|       | 9.1.2.2.1.2      | PS::DomainInfo::initialize   | 107    |
|       | 9.1.2.2.1.3      | PS:: Domain Info:: set Number Of Domain Multi Dimensional Set of the Set of S | on 108 |
|       | 9.1.2.2.1.4      | PS::DomainInfo::setBoundaryCondition   | 108    |
|       | 9.1.2.2.1.5      | PS::DomainInfo::getBoundaryCondition   | 109    |
|       | 9.1.2.2.1.6      | PS::DomainInfo::setPosRootDomain   | 109    |
|       | 9.1.2.2.2        | Decomposition of Domain  | 109    |
|       | 9.1.2.2.2.1      | PS::DomainInfo::collectSampleParticle  | 110    |
|       | 9.1.2.2.2.2      | PS::DomainInfo::decomposeDomain  | 111    |
|       | 9.1.2.2.2.3      | PS::DomainInfo::decomposeDomainAll   | 112    |
|       | 9.1.2.2.3        | Time Measurment  | 113    |
|       | 9.1.2.2.3.1      | PS::DomainInfo::getTimeProfile   | 113    |
|       |                  | PS::DomainInfo::clearTimeProfile   | 113    |
|       | 9.1.2.2.4        | Obtain Information   | 114    |
|       | 9.1.2.2.4.1      | PS::DomainInfo::getMemSizeUsed   | 114    |
| 9.1.3 | ParticleSystem C | lass   | 114    |
|       | 9.1.3.1 Creation | n of Object  | 114    |
|       | 9.1.3.2 API      |  | 115    |
|       | 9.1.3.2.1        | Initial Setup  | 115    |
|       | 9.1.3.2.1.1      | Constructor  | 115    |
|       | 9.1.3.2.1.2      | PS::ParticleSystem::initialize   | 115    |
|       | 9.1.3.2.1.3      | PS::ParticleSystem::   |        |
|       |                  | set Average Target Number Of Sample Particle Per Processor (Appendix Per Pro | ss116  |
|       | 9.1.3.2.2        | Obtain Information   | 116    |
|       | 9.1.3.2.2.1      | PS::ParticleSystem::operator []  | 117    |
|       | 9.1.3.2.2.2      | PS:: Particle System:: get Number Of Particle Local  .  .  | 117    |
|       | 9.1.3.2.2.3      | $PS:: Particle System:: get Number Of Particle Global \ . \ .$   | 117    |
|       | 9.1.3.2.2.4      | PS::DomainInfo::getUsedMemorySize  | 118    |
|       | 9.1.3.2.3        | File I/O   | 118    |
|       | 9.1.3.2.3.1      | PS::ParticleSystem::readParticleAscii  | 121    |
|       | 9.1.3.2.3.2      | $PS:: Particle System:: read Particle Binary \ . \ . \ . \ . \ . \ .$  | 123    |
|       | 9.1.3.2.3.3      | PS::ParticleSystem::writeParticleAscii   | 125    |
|       | 9.1.3.2.3.4      | PS::ParticleSystem::writeParticleBinary  | 127    |
|       | 9.1.3.2.4        | Exchange Particles   | 128    |
|       | 9.1.3.2.4.1      | PS::ParticleSystem::exchangeParticle   | 129    |
|       | 9.1.3.2.5        | Adding and removing particles  | 129    |
|       | 9.1.3.2.5.1      | $PS:: Particle System:: add One Particle () \ . \ . \ . \ . \ . \ .$   | 129    |
|       | 9.1.3.2.5.2      | $PS:: Particle System:: remove Particle() \ \dots \ \dots \ \dots$   | 129    |
|       | 9.1.3.2.6        | Time Measurement   | 130    |
|       | 9.1.3.2.6.1      | $PS:: Particle System:: get Time Profile \ . \ . \ . \ . \ . \ . \ .$  | 130    |
|       | 9.1.3.2.6.2      | $PS:: Particle System:: clear Time Profile \ . \ . \ . \ . \ .$  | 131    |
|       | 9.1.3.2.7        | Others   | 131    |
|       | 9.1.3.2.7.1      | PS:: Particle System:: adjust Position Into Root Domain  | 131    |

|       | 9.1.3.2.7.2        | PS::ParticleSystem::setNumberOfParticleLocal   | 132 |
|-------|--------------------|--|-----|
|       | 9.1.3.2.7.3        | PS::ParticleSystem::sortParticle   | 132 |
| 9.1.4 | TreeForForce Class | SS   | 133 |
|       | 9.1.4.1 Creation   | n of Object  |     |
|       | 9.1.4.1.1          |  |     |
|       | 9.1.4.1.2          | PS::SEARCH_MODE_LONG_SCATTER   | 134 |
|       | 9.1.4.1.3          | PS::SEARCH_MODE_LONG_SYMMETRY  | 135 |
|       | 9.1.4.1.4          | PS::SEARCH_MODE_LONG_CUTOFF  | 135 |
|       | 9.1.4.1.5          | PS::SEARCH_MODE_GATHER   | 136 |
|       | 9.1.4.1.6          | PS::SEARCH_MODE_SCATTER  | 136 |
|       | 9.1.4.1.7          | PS::SEARCH_MODE_SYMMETRY   | 136 |
|       | 9.1.4.2 API        |  | 137 |
|       | 9.1.4.2.1          | Initial Setup  | 137 |
|       | 9.1.4.2.1.1        | Constructor  | 138 |
|       | 9.1.4.2.1.2        | PS::TreeForForce::initialize   | 138 |
|       | 9.1.4.2.2          | Low Level APIs   | 139 |
|       | 9.1.4.2.2.1        | PS::TreeForForce::setParticleLocalTree   | 140 |
|       | 9.1.4.2.2.2        | PS::TreeForForce::makeLocalTree  | 140 |
|       | 9.1.4.2.2.3        | PS::TreeForForce::makeGlobalTree   | 141 |
|       | 9.1.4.2.2.4        | PS::TreeForForce::calcMomentGlobalTree   | 141 |
|       | 9.1.4.2.2.5        | PS::TreeForForce::calcForce  | 142 |
|       | 9.1.4.2.2.6        | PS::TreeForForce::getForce   | 143 |
|       |                    | High Level APIs  |     |
|       |                    | PS::TreeForForce::calcForceAllAndWriteBack   |     |
|       | 9.1.4.2.3.2        | PS::TreeForForce::calcForceAll   | 151 |
|       |                    | PS::TreeForForce::calcForceAndWriteBack  |     |
|       | 9.1.4.2.4          | Neighbor List  | 155 |
|       | 9.1.4.2.4.1        | getNeighborListOneParticle   |     |
|       | 9.1.4.2.5          |  |     |
|       | 9.1.4.2.5.1        | PS::TreeForForce::getTimeProfile   |     |
|       |                    | PS::TreeForForce::clearTimeProfile   |     |
|       |                    | Obtain Information   |     |
|       |                    | PS::TreeForForce::getNumberOfInteractionEPEPLoc  |     |
|       |                    | PS::TreeForForce::getNumberOfInteractionEPEPGlo  |     |
|       |                    | ${\bf PS::} {\bf TreeForForce::} {\bf getNumberOfInteractionEPSPLoc}$  |     |
|       |                    | PS:: Tree For Force:: get Number Of Interaction EPSPG location and the property of the prope |     |
|       |                    | PS::TreeForForce::clearNumberOfInteraction   | 159 |
|       |                    | PS::TreeForForce::getNumberOfWalkLocal   | 159 |
|       |                    | PS::TreeForForce::getNumberOfWalkGlobal  | 160 |
|       |                    | PS::TreeForForce::getMemSizeUsed   | 160 |
|       | 9.1.4.2.7          |  | 160 |
|       |                    | getEpjFromId   | 160 |
| 9.1.5 | Comm Class         |  | 161 |
| -     |                    |  | 161 |
|       | 9.1.5.1.1          | PS::Comm::getRank  | 162 |
|       |                    | PS::Comm::getNumberOfProc  | 162 |

|           |      |        | 9.1.5.1.3 PS::Comm::getRankMultiDim                                  | 162 |
|-----------|------|--------|--|-----|
|           |      |        | 9.1.5.1.4 PS::Comm::getNumberOfProcMultiDim                          | 163 |
|           |      |        | $9.1.5.1.5  {\rm PS::Comm::synchronizeConditionalBranchAND}  .  .$   | 163 |
|           |      |        | 9.1.5.1.6 PS::Comm::synchronizeConditionalBranchOR                   | 163 |
|           |      |        | 9.1.5.1.7 PS::Comm::getMinValue                                      | 164 |
|           |      |        | 9.1.5.1.8 PS::Comm::getMaxValue                                      | 164 |
|           |      |        | 9.1.5.1.9 PS::Comm::getSum   | 165 |
|           |      |        | 9.1.5.1.10 PS::Comm::broadcast                                       | 165 |
|           |      | 9.1.6  | Other functions  | 166 |
|           |      | 0.1.0  | 9.1.6.1 Timing function  | 166 |
|           |      |        | 9.1.6.1.1 PS::GetWtime   | 166 |
|           | 9.2  | Extend | led Classes  | 166 |
|           | 5.2  | 9.2.1  | Summary  | 166 |
|           |      | 9.2.2  | Particle Mesh Class  | 166 |
|           |      | 9.4.4  | 9.2.2.1 Creation of Object   | 166 |
|           |      |        | ů  | 167 |
|           |      |        |  |     |
|           |      |        | 9.2.2.2.1 Initial Setup  | 167 |
|           |      |        | 9.2.2.2.1.1 Constructor  | 167 |
|           |      |        | 9.2.2.2.2 Low Level API  | 167 |
|           |      |        | 9.2.2.2.2.1 PS::PM::ParticleMesh::setDomainInfoParticleMesh          | 168 |
|           |      |        | 9.2.2.2.2.2 PS::PM::ParticleMesh::setParticleParticleMesh            | 168 |
|           |      |        | 9.2.2.2.3 PS::PM::ParticleMesh::calcMeshForceOnly                    | 169 |
|           |      |        | 9.2.2.2.2.4 PS::PM::ParticleMesh::getForce                           | 169 |
|           |      |        | 9.2.2.2.5 PS::PM::ParticleMesh::getPotential                         | 170 |
|           |      |        | 9.2.2.2.3 High Level API   | 170 |
|           |      |        | $9.2.2.2.3.1 \ PS::PM::ParticleMesh::calcForceAllAndWriteBack \ .$   | 170 |
|           |      |        | 9.2.2.3 Predefined Macros  | 171 |
|           |      |        | 9.2.2.4 How To Use Particle Mesh Class                               | 172 |
|           |      |        | 9.2.2.4.1 Compile of Particle Mesh Class                             | 173 |
|           |      |        | 9.2.2.4.2 Writing Source Code  | 173 |
|           |      |        | 9.2.2.4.3 Compile of Source Code                                     | 173 |
|           |      |        | 9.2.2.4.4 Note   |     |
|           |      | 9.2.3  | Phantom-GRAPE for X86  |     |
|           |      |        |  |     |
| <b>10</b> | Dete | ection | of Errors  | 174 |
|           |      |        | et   | 174 |
|           | 10.2 | Compi  | le time errors   | 174 |
|           | 10.3 | Run ti | me errors  | 174 |
|           |      | 10.3.1 | PS_ERROR: can not open input file                                    | 174 |
|           |      | 10.3.2 | PS_ERROR: can not open output file                                   | 175 |
|           |      |        | PS_ERROR: Do not initialize the tree twice                           | 175 |
|           |      |        | PS_ERROR: The opening criterion of the tree must be $>=0.0\ldots$    | 175 |
|           |      |        | PS_ERROR: The limit number of the particles in the leaf cell must be |     |
|           |      |        | > 0  | 175 |
|           |      | 10.3.6 | PS_ERROR: The limit number of particles in ip groups msut be >=      |     |
|           |      |        | that in leaf cells   | 175 |
|           |      |        |  |     |

|                  | 10.3.7   | FDPS limit number   | 176        |
|------------------|----------|---|------------|
|                  | 10.3.8   | PS_ERROR: The forces w/o cutoff can be evaluated only under the                                 | _, ,       |
|                  |          | open boundary condition   | 176        |
|                  |          | PS_ERROR: A particle is out of root domain  | 176        |
|                  |          | is must between 0 and 1   | 176        |
|                  |          | 1 PS_ERROR: The coodinate of the root domain is inconsistent 2 PS_ERROR: Vector invalid accesse | 177<br>177 |
| 11 Kn            | own B    | ngs   | 178        |
| 12 Lir           | nitatior | 1   | 179        |
| $13~\mathrm{Us}$ | er Supp  | port  | 180        |
| 14 Lic           | ense     |   | 181        |
| 15 Ch            | ange L   | $\mathbf{p}$  | 182        |
| A Ex             | amples   | of implementation using use-defined classes   | 185        |
| A.1              | Class    | FullParticle  | 185        |
|                  | A.1.1    | Summary   | 185        |
|                  | A.1.2    | Presumptions  | 185        |
|                  | A.1.3    | Necessary member functions  | 185        |
|                  |          | A.1.3.1 Summary   | 185        |
|                  |          | A.1.3.2 FP::getPos  | 185        |
|                  |          | A.1.3.3 FP::copyFromForce   | 186        |
|                  | A.1.4    | Member functions necessary for some situations  | 186        |
|                  |          | A.1.4.1 Summary   | 186        |
|                  |          | A.1.4.2 The cases where all the types of PS::SEARCH_MODE but type                               |            |
|                  |          | PS::SEARCH_MODE_LONG are used   | 187        |
|                  |          | A.1.4.2.1 FP::getRSearch  | 187        |
|                  |          | A.1.4.3 The case where file I/O APIs of class ParticleSystem is used                            | 187        |
|                  |          | A.1.4.3.1 FP::readAscii   | 188        |
|                  |          | A.1.4.3.2 FP::writeAscii  | 188        |
|                  |          | A.1.4.3.3 FP::readBinary  | 189        |
|                  |          | A.1.4.3.4 FP::writeBinary   | 190        |
|                  |          | A.1.4.4 The case where ParticleSystem::adjustPositionIntoRootDomais                             | n          |
|                  |          | is used   | 191        |
|                  |          | A.1.4.4.1 FP::setPos  | 191        |
|                  |          | A.1.4.5 The case where class Particle Mesh is used  | 191        |
|                  |          | A.1.4.5.1 FP::getChargeParticleMesh   | 191        |
|                  |          | A.1.4.5.2 FP::copyFromForceParticleMesh   | 192        |
| A.2              | 2 Class  | EssentialParticleI  | 193        |
|                  | A.2.1    | Summary   | 193        |

|     | A.2.2  | Presumptions  | 93         |
|-----|--------|---|------------|
|     | A.2.3  | Necessary member functions                                  | 93         |
|     |        |   | 93         |
|     |        |   | 93         |
|     |        |   | 94         |
|     | A.2.4  |   | 95         |
|     |        | · · · · · · · · · · · · · · · · · · ·                       | 95         |
|     |        | A.2.4.2 The case where PS::SEARCH_MODE_GATHER and           |            |
|     |        | PS::SEARCH_MODE_SYMMETRY are adopted for type               |            |
|     |        | <u>* -</u>  | 95         |
|     |        |   | 95         |
| A.3 | Class  | 9   | 95         |
|     | A.3.1  |   | 95         |
|     | A.3.2  | · ·   | 96         |
|     | A.3.3  | 1   | 96         |
|     |        | y .   | 96         |
|     |        | v   | 96         |
|     |        | 8   | 97         |
|     | A.3.4  | 1 /   | 98         |
|     | 11.0.1 |   | .98        |
|     |        | A.3.4.2 The cases where all but PS::SEARCH_MODE_LONG are    |            |
|     |        |   | 98         |
|     |        |   | .98        |
|     |        | A.3.4.3 The cases where all but PS::BOUNDARY_CONDITION_OPEN |            |
|     |        |   | 99         |
|     |        |   | 99         |
| Δ 1 | Class  |   | 99         |
| л.4 | A.4.1  |   | 99         |
|     | A.4.2  |   | 200        |
|     | A.4.2  |   | 200        |
|     |        | v   | 200        |
|     |        |   | 200        |
|     |        | -   | 200        |
|     |        | <b>V</b>  | 201        |
|     |        | 1   | .01<br>202 |
|     |        | <u>.</u>  | $02 \\ 02$ |
|     |        | v i   |            |
|     |        |   | 203        |
|     | 1 1 2  | <u> </u>  | 203        |
|     | A.4.3  | v   | 204        |
|     |        | v   | 204        |
|     |        |   | 204        |
|     |        |   | 205        |
|     |        | 8   | 206        |
|     |        |   | 206        |
|     |        | A.4.3.6 Mom::accumulateAtLeaf                               | 07         |
|     |        | A 4.2.7 Mamusaaymulata                                      | 00         |

|      |         | A.4.3.8 Mom::set                               |     |
|------|---------|--|-----|
|      |         | A.4.3.9 Mom::accumulateAtLeaf2                 | 210 |
|      |         | A.4.3.10 Mom::accumulate2                      | 211 |
| A.5  | Class S | SuperParticleJ                                 | 212 |
|      | A.5.1   | Summary  |     |
|      | A.5.2   |  | 212 |
|      |         |  | 212 |
|      |         |  | 212 |
|      |         | A.5.2.1.2 PS::SPJQuadrupole                    |     |
|      |         | A.5.2.1.3 PS::SPJMonopoleGeometricCenter       |     |
|      |         | A.5.2.1.4 PS::SPJDipoleGeometricCenter         |     |
|      |         | A.5.2.1.5 PS::SPJQuadrupoleGeometricCenter     |     |
|      |         | A.5.2.2 PS::SEARCH_MODE_LONG_CUTOFF            |     |
|      |         |  | 215 |
|      | A.5.3   |  | 215 |
|      |         | v  | 215 |
|      |         |  | 216 |
|      |         |  | 216 |
|      |         | A.5.3.4 SPJ::copyFromMoment                    |     |
|      |         |  | 218 |
|      |         |  | 219 |
| A.6  | Class 1 |  | 219 |
|      | A.6.1   |  | 219 |
|      | A.6.2   | v  | 219 |
|      | A.6.3   | 1  | 219 |
|      |         | v  | 220 |
| A.7  | Class 1 |  | 220 |
|      | A.7.1   |  | 220 |
|      |         | ·  | 220 |
|      | A.7.3   | Member functions necessary for some situations |     |
|      |         | A.7.3.1 Hdr::readAscii                         |     |
|      |         | A.7.3.2 Hdr::writeAscii                        |     |
| A.8  | Functi  |  | 222 |
|      | A.8.1   |  | 222 |
|      | A.8.2   | ·  | 222 |
|      | A.8.3   | 1  | 222 |
| A.9  |         | 9  | 224 |
|      | A.9.1   |  | 224 |
|      | A.9.2   | v  | 224 |
|      | A.9.3   | -  | 224 |
| A.10 |         |  | 226 |
|      |         |  | 226 |
|      |         | v  | 226 |
|      |         |  | 226 |
| A.11 |         | 1  | 230 |
|      |         |  | 230 |

| A.11.2 Pre | mises . |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  |  | 23 | 30 | ) |
|------------|---------|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|----|----|---|
|------------|---------|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|--|----|----|---|

# 1 Concept of Document

This document is the specification of FDPS (Framework for Developing Particle Simulator), which supports the development of massively parallel particle simulation codes. This document is written by Ataru Tanikawa, Masaki Iwasawa, Natsuki Hosono, Keigo Nitadori, Takayuki Muranushi, and Junichiro Makino at RIKEN Advanced Institute for Computational Science.

This document is structured as follows.

In sections 2, 3, and 4, we present prerequisites for programing with FDPS. In section 2, we show the concept of FDPS. In section 3, we present the file configuration of FDPS. In section 4, we describe how to compile simulation codes with FDPS.

In sections 5, 6, 7, 8, and 9, we present necessary information to develop simulation codes with FDPS. In section 5, we describe the structure of namespaces unused in FDPS. In section 6, we described data types defined in FDPS. In section 7, we introduce user-defined classes and function objects necessary for developing codes with FDPS. In section 8, we describe APIs used to initialize and finalize FDPS. In section 9, we present modules in FDPS and their APIs.

In sections 10, 11, 12, and 13, we present information useful for troubleshooting. In section 10, we describe error messages. In section 11, we present known bugs. In section 12, we describe the limitation of FDPS. In section 13, we present our current system for user supports.

Finally, we describe the license of FDPS in section 14, and the change log of this document in section 15.

# 2 Concept of FDPS

In this section, we present the design concept of FDPS: the purpose of its development, the basic concept, and the behavior of codes developed using FDPS.

# 2.1 Purpose of the development of FDPS

In the fields of science and engineering, particle method is used for a wide variety of simulations, such as gravitational N-body simulation, Smoothed Particle Hydrodynamics (SPH) simulation, vortex method, Moving Particle Semi-implicit (MPS) method, molecular dynamics simulation, and so on. We need high-performance particle simulation codes in order to follow physical phenomena with high spatial resolution, and for long timescales.

We cannot avoid parallelization in order to develop high-performance particle simulation codes. For the parallelization, we need to implement the following procedures: dynamic domain decomposition for load balancing, exchange of particles between computing nodes, optimization of communication among nodes, effective use of cache memories and SIMD operation units, and support for accelerators. So far, individual research groups were trying to implement these procedures.

However, the above procedures are necessary for any particle simulation codes. The purpose of the development of FDPS is to provide numerical libraries for implementing these procedures, and reduce researchers' and programmers' burdens. We will be happy if researchers and programmers can use their time more creatively by making use of FDPS.

# 2.2 Basic concept

In this section, we describe the basic concept of FDPS.

#### 2.2.1 Procedures of massively parallel particle simulations

First, we describe our model of massively parallel particle simulations on FDPS. In particle simulations, the set of ordinary differential equations,

$$\frac{d\mathbf{u}_i}{dt} = \sum_i f(\mathbf{u}_i, \mathbf{u}_j) + \sum_s g(\mathbf{u}_i, \mathbf{v}_s), \tag{1}$$

is numerically integrated, where  $u_i$  is the quantity vector of *i*-particle. This vector includes quantities of particle *i*, such as mass, position, and velocity. The function f specifies a force exerted by particle j on particle i. Hereafter, a particle receiving a force is called i-particle, and a particle exerting a force is called particle j. The vector  $v_s$  is the quantity vector of a superparticle which is the representative particle of a group of particles distant from i-particle. The function g specifies a force exerted by a superparticle on a particle. The second term in the left hand side of eq. (1) is non-zero in the case of long-range forces (e.g. gravity and Coulomb force), while it is zero in the case of short-range forces (e.g. pressure of fluid).

Massively parallel simulation codes integrate the above eq. (1) in the following steps (initialization and data I/O are omitted).

1. In the following two steps, we determine which MPI process handles which particles.

- (a) Decompose the whole domain into subdomains, and determine which MPI process handles which subdomains, in order to balance the calculation cost (domain decomposition).
- (b) MPI processes exchange their particles in order for each MPI process to have particles in its subdomain.
- 2. Each MPI process gathers quantity vectors of j-particles  $(\boldsymbol{u}_j)$  and superparticles  $(\boldsymbol{v}_s)$  required to calculate forces exerted on i-th i-particle (making interaction lists).
- 3. Each MPI process calculates the right hand of eq. (1) for all of its *i*-particle and obtains  $d\mathbf{u}_i/dt$ .
- 4. Each MPI process performes the time integration of its *i*-particles by using quantity vectors of  $\mathbf{u}_i$  and their derivatives  $d\mathbf{u}_i/dt$ .
- 5. Return to step 1.

# 2.2.2 Division of tasks between users and FDPS

FDPS handles tasks related to interaction calculation and its efficient parallelization and the user-written code performs the rest. The actual function for interaction calculation is supplied by users. Thus, FDPS deals with domain decomposition and exchange of particles (step 1), and making interaction lists. On the other hand, the user code is responsible for actual calculation of forces (step 3), and time integration (step 4). Users can avoid the development of complicated codes necessary to realize massively parallel program, by utilizing FDPS APIs.

#### 2.2.3 Users' tasks

Users's tasks are as follows.

- Define a particle (section 7). Users need to specify quantities of particles, *i.e.* the quantity vector  $u_i$  in eq. (1), which contains quantities such as position, velocity, acceleration, chemical composition, and particle size.
- Define interaction (section 7). Users need to specify the interaction between particles, *i.e.* the function f and g in eq. (1), such as gravity, Coulomb force, and pressure.
- Call FDPS APIs (setion 8 and 9).
- Time integration of particles, diagnostic, output etc.

# 2.2.4 Complement

The right hand side of eq. (1), the particle-particle interactions, is strictly of two-body nature. FDPS APIs can not be used to implement three-particle interactions. However, for example, FDPS has APIs to return neighbor lists. Users can calculate three- or more-body interactions, using these neighbor lists.

Calculation steps in section 2.2.1 imply that all particles have one same timestep. FDPS APIs do not support individual timestep scheme. However, users can develop a particle simulation code with individual timestep scheme, using the Particle Particle Particle Tree method.

# 2.3 The structure of a simulation code with FDPS

We overview the structure of a simple simulation code written using FDPS. In a code with FDPS, three FDPS-supplied classes and several user-defined classes are used.

- DomainInfo class. This class contains the information of all the subdomains, and APIs for domain decomposition.
- ParticleSystem class. This class contains the information of all particles in each MPI process, and APIs for the exchange of particles among MPI processes.
- TreeForForce class. This class contains tree structure made from particle distribution, and APIs for making interaction lists.
- User-defined classes. These classes include the definitions of particles and interactions.

These classes communicate with each other. This is illustrated in fig. 1. The communication in this figure corresponds to steps 1 and 2, and to initialization (step 0).

- 0. Users give a user-defined particle class to ParticleSystem class, and a function object to TreeForForce class. These are not class inheritance. The particle class is used as a template argument of ParticleSystem class, and the function object is used as an argument of APIs in TreeForForce class.
- 1. Do load balancing in the following two steps.
  - (a) The user code calls APIs for domain decomposition in DomainInfo class. Particle information is transfered from ParticleSystem class to DomainInfo class (red text and arrows).
  - (b) The user code calls APIs for exchange of particles in ParticleSystem class. Information of subdomains is transferred from DomainInfo class to ParticleSystem class (blue text and arrows).
- 2. Do the force calculation in the following steps.
- (a) The user code calls force calculation API.
- (b) FDPS makes interaction lists in TreeForForce class. Information of subdomains and particles is transferd from DomainInfo and ParticleSystem classes (green text and arrows).
- (c) FDPS calls an user-defined function object. This API is included in TreeForForce class. Interactions are calculated, and the results are transferred from TreeForForce class to ParticleSystem class (gray text and arrows).

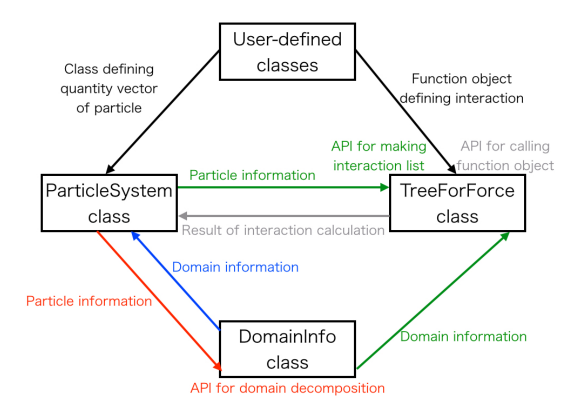


Figure 1: Illustration of module interface and data transfer.

# 3 Files and directories

# 3.1 Summary

We describe the directory strutcures and files of FDPS file distribution.

#### 3.2 Documents

Document files are in directory doc. Files doc\_tutorial\_e.pdf and doc\_specs\_en.pdf are tutorial and specification, respectively.

#### 3.3 Source files

Source files are in directory src. Files related to the standard features are directly under directory src. When users include header file particle\_simulator.hpp in their source files, the standard features become available.

#### 3.3.1 Extended features

Source files related to extended features are in directories in directory src. The extended features include Particle Mesh and Phantom-GRAPE version x86.

#### 3.3.1.1 Particle Mesh scheme

Source files related to the Particle Mesh scheme are in directory particle\_mesh. When users edit Makefile, and run make, they get header file particle\_mesh\_class.hpp and library libpm.a. By including this header file, and linking this library, they can use the features of the Particle Mesh scheme.

#### 3.3.1.2 x86 version Phantom-GRAPE

Source files related to x86 version Phantom-GRAPE are in directory src/phantom\_GRAPE\_x86. This directory has directories containing Phantom-GRAPE libraries for low precision, for low precision with cutoff, and for high precision.

#### 3.3.1.2.1 Low precision

Files are in directory src/phantom\_GRAPE\_x86/G5/newton/libpg5. Users should edit Makefile in this directory, and run make, to get library libpg5.a. Users include header file gp5util.h in this directory, and to link library libpg5.a, to use this Phantom-GRAPE.

#### 3.3.1.2.2 Low precision with cutoff

Files are in directory src/phantom\_GRAPE\_x86/G5/table. Users should edit Makefile in this directory, and run make, to get library libpg5.a. Users include header file gp5util.h in this directory, and to link library libpg5.a, to use this Phantom-GRAPE.

#### 3.3.1.2.3 High precision

Files are in directory src/phantom\_GRAPE\_x86/G6/libavx. Users should edit Makefile in this directory, and run make, to get library libg6avx.a. Users include header file gp6util.h in this directory, and to link library libg6avx.a, to use this Phantom-GRAPE.

#### 3.4 Test codes

Test codes are in directory tests. When users run make check in this directory, the test suite are executed.

# 3.5 Sample codes

Sample codes are in directory sample. There are two sample codes: gravitational N-body simulation, and SPH simulation.

# 3.5.1 Gravitational N-body simulation

Source files are in directory sample/nbody. In file doc\_tutorial.pdf, we describe how to use this code.

# 3.5.2 SPH Simulation

Source files are in directory sample/sph. In file doc\_tutorial.pdf, we describe how to use this code.

# 4 Macros at compile time

# 4.1 Summary

Some of the features of FDPS should be specified at the compile time. They are (a)coordinate system, (b)method of parallelization, and (c) accuracy of floating point types.

# 4.2 Coordinate system

# 4.2.1 Summary

Users have alternatives of 2D and 3D Cartesian coordinate systems.

# 4.2.2 3D Cartesian coordinate system

3D Cartesian coordinate system is used by default.

### 4.2.3 2D Cartesian coordinate system

2D Cartesian coordinate system can be used by defining PARTICLE\_SIMULATOR\_TWO\_DIMENSION as macro.

# 4.3 Parallel processing

# 4.3.1 Summary

Users choose whether OpenMP is used or not, and whether MPI is used or not.

# 4.3.2 OpenMP

OpenMP is disabled by default. If macro PARTICLE\_SIMULATOR\_THREAD\_PARALLEL is defined, OpenMP becomes enabled. Compiler option "-fopenmp" is required for GCC compiler.

#### 4.3.3 MPI

MPI is disabled by default. If macro PARTICLE\_SIMULATOR\_MPI\_PARALLEL is defined.

# 4.4 Accuracy of data types

# 4.4.1 Summary

Users can specify the accuracy of data types of Moment classes described in § 7.5.2) and SuperParticleJ classes described in § 7.6.2.

#### 4.4.2 Accuracy of SuperParticleJ and Moment classes prepared in FDPS

All the member variables in SuperParticleJ classes and Moment classes are 64 bit accuracy by default. They becomes 32 bit accuracy if macro PARTICLE\_SIMULATOR\_SPMOM\_F32BIT is defined at the compile time.

# 5 Namespace

# 5.1 Summary

In this section, we describe the namespaces used in FDPS. All the FDPS APIs are under namespace ParticleSimulator. In the following, we show APIs directly under ParticleSimulator, and namespaces nested under ParticleSimulator.

# 5.2 ParticleSimulator

The standard features of FDPS are in namespace ParticleSimulator.

Namespace ParticleSimulator is abbreviated to PS as follows.

```
namespace PS = ParticleSimulator;
```

In this document, we use this abbreviation.

Extended features of FDPS are grouped under nested namespaces. Currently, ParticleMesh features is under nested namespace ParticleMesh. In the following, we describe this namespace containing the extended features.

#### 5.2.1 ParticleMesh

The features of Particle Mesh are under namespace PartcielMesh. This namespace is abbreviated to PM as follows.

```
ParticleSimulator {
    ParticleMesh {
    }
    namespace PM = ParticleMesh;
}
```

In this document, we use this abbreviation.

# 6 Data Type

# 6.1 Summary

In this chapter we define data types used in FDPS. The data types include integer types, floating point types, vector types, symmetric matrix types, PS::SEARCH\_MODE type, and enumerated types. We recommend users to use these data types. The integer and floating point types can be replaced with data types available in C/C++ languages. PS::SEARCH\_MODE and enumerated types must be used. In the following, we describe these data types.

# 6.2 Integer type

#### 6.2.1 Overview

The integer types are PS::S32, PS::S64, PS::U32, and PS::U64. We describe these data types in this section.

#### 6.2.2 PS::S32

PS::S32, which is 32-bit signed integer, is defined as follows.

```
Listing 1: S32
```

```
1 namespace ParticleSimulator {
2    typedef int S32;
3 }
```

#### 6.2.3 PS::S64

PS::S64, which is 64-bit signed integer, is defined as follows.

```
Listing 2: S64
```

```
1 namespace ParticleSimulator {
2     typedef long long int S64;
3 }
```

### 6.2.4 PS::U32

PS::U32, which is 32-bit unsigned integer, is defined as follows.

```
Listing 3: U32
```

```
namespace ParticleSimulator {
2 typedef unsigned int U32;
3 }
```

#### 6.2.5 PS::U64

PS::U64, which is 64-bit unsigned integer, is defined as follows.

```
Listing 4: U64
```

```
1 namespace ParticleSimulator {
2     typedef unsigned long long int U64;
3 }
```

# 6.3 Floating point type

#### 6.3.1 Abstract

The floating point types are PS::F32 and PS::F64. We described these data types in this section.

#### 6.3.2 PS::F32

PS::F32, which is 32-bit floating point number, is defined as follows.

#### Listing 5: F32

```
1 namespace ParticleSimulator {
2     typedef float F32;
3 }
```

#### 6.3.3 PS::F64

PS::F64, which is 64-bit floating point number, is defined as follows.

# Listing 6: F64

```
1 namespace ParticleSimulator {
2    typedef double F64;
3 }
```

# 6.4 Vector type

#### 6.4.1 Abstract

The vector types are PS::Vector2 (2D vector) and PS::Vector3 (3D vector). We describe these vector types first. These are template classes, which can take basic datatypes as F32 or F64 as template arguments. We then present wrappers for these vector types.

#### 6.4.2 PS::Vector2

PS::Vector2 has two components: x and y. We define various APIs and operators for these components. In the following, we list them.

```
1 namespace ParticleSimulator{
2
       template <typename T>
3
       class Vector2{
4
       public:
           // Two member variables
5
6
           T x, y;
7
8
           // Constructors
9
           Vector2();
10
           Vector2(const T _x, const T _y) : x(_x), y(_y) {}
           Vector2(const T s) : x(s), y(s) {}
11
12
           Vector2(const Vector2 & src) : x(src.x), y(src.y) {}
13
14
           // Assignment operator
15
           const Vector2 & operator = (const Vector2 & rhs);
16
17
           //[] operators
18
           const T & opertor[](const int i);
           T & operator[](const int i);
19
20
21
           // Addition and subtraction
           Vector2 operator + (const Vector2 & rhs) const;
22
23
           const Vector2 & operator += (const Vector2 & rhs);
24
           Vector2 operator - (const Vector2 & rhs) const;
25
           const Vector2 & operator -= (const Vector2 & rhs);
26
27
           // Scalar multiplication
28
           Vector2 operator * (const T s) const;
29
           const Vector2 & operator *= (const T s);
30
           friend Vector2 operator * (const T s,
31
                                        const Vector2 & v);
32
           Vector2 operator / (const T s) const;
33
           const Vector2 & operator /= (const T s);
34
35
           // Inner product (scalar product)
36
           T operator * (const Vector2 & rhs) const;
37
           // Outer product (vector product).
38
39
           // Caution: the return value is a scalar value.
40
           T operator ^ (const Vector2 & rhs) const;
41
42
           // Typecast to Vector2<U>
43
           template <typename U>
44
           operator Vector2<U> () const;
45
       };
```

```
46 }
47 namespace PS = ParticleSimulator;
```

#### 6.4.2.1 Constructor

```
template<typename T>
PS::Vector2<T>::Vector2()
```

# • Argument

None.

#### • Feature

Default constructor. The member variables  $\mathbf{x}$  and  $\mathbf{y}$  are initialized as 0.

```
template<typename T>
PS::Vector2<T>::Vector2(const T_x, const T_y)
```

### • Argument

```
_x: Input. Type const T.
_y: Input. Type const T.
```

### • Feature

Values of members x and y are set to values of arguments \_x and \_y, respectively.

```
template<typename T>
PS::Vector2<T>::Vector2(const T s);
```

#### • Argument

s: Input. Type const T.

#### • Feature

Values of both of members x and y are set to the value of argument s.

# 6.4.2.2 Copy constructor

```
template<typename T>
PS::Vector2<T>::Vector2(const PS::Vector2<T> & src)
```

### • Argument

src: Input. Type const PS::Vector2<T>.

#### • Feature

Copy constructor. The new variable will have the same value as src.

#### 6.4.2.3 Member variables

```
template<typename T>
T PS::Vector2<T>::x;

template<typename T>
T PS::Vector2<T>::y;
```

#### • function

Member variables, x and y can be directly handled.

#### 6.4.2.4 Assignment operator

### • Argument

rhs: Input. Type const PS::Vector2<T> &.

#### • Return value

Type const PS::Vector2<T> &. Assigns the values of components of rhs to its components, and returns the vector itself.

# 6.4.2.5 [] operators

#### • Argument

i: input. Type const int.

#### • Return value

Type const < T > &. It returns i-component of the vector.

#### • Remark

This operator would be slower than the direct access of the member variables.

#### • Argument

i: input. Type const int.

#### • Return value

Type  $\langle T \rangle$  &. It returns i-component of the vector.

#### • Remark

This operator would be slower than the direct access of the member variables.

#### 6.4.2.6 Addition and subtraction

#### • Argument

rhs: Input. Type const PS::Vector2<T> &.

#### • Return value

Type PS::Vector2 < T >. Add the components of rhs and its components, and return the results.

#### • Argument

rhs: Input. Type const PS::Vector2<T> &.

#### • Return value

Type const PS::Vector2 < T > &. Add the components of rhs to its components, and return itself (lhs is changed).

# • Argument

rhs: Input. Type const PS::Vector2<T> &.

#### • Return value

Type PS::Vector2 < T >. Subtract the components of rhs from its components, and return the results.

### • Argument

rhs: Input. Type const PS::Vector2<T> &.

#### • Return value

Type const PS::Vector2 < T > &. Subtract the components of rhs from its components, and return itself (lhs is changed).

# 6.4.2.7 Scalar multiplication

```
template<typename T>
PS::Vector2<T> PS::Vector2<T>::operator * (const T s) const;
```

#### • Argument

s: Input. Type const T.

#### • Return value

Type PS::Vector2<T>. Multiply its components by s, and return the results.

```
template<typename T>
const PS::Vector2<T> & PS::Vector2<T>::operator *= (const T s);
```

### • Argument

rhs: Input. Type const T.

#### • Return value

Type const PS::Vector2 < T > &. Multiply its components by s, and return itself (lhs is changed).

```
template<typename T>
PS::Vector2<T> PS::Vector2<T>::operator / (const T s) const;
```

#### • Argument

s: Input. Type const T.

#### • Return value

Type PS::Vector2 < T >. Divide its components by s, and return the results.

```
template<typename T>
const PS::Vector2<T> & PS::Vector2<T>::operator /= (const T s);
```

#### • Argument

rhs: Input. Type const T.

#### • Return value

Type const PS::Vector2 < T > &. Divide its components by s, and return itself (lhs is changed).

### 6.4.2.8 Inner and outer products

```
template<typename T>
T PS::Vector2<T>::operator * (const PS::Vector2<T> & rhs) const;
```

# • Argument

rhs: Input. Type const PS::Vector2<T> &.

#### • Return value

Type T. Take inner product of it and rhs, and return the result.

```
template<typename T>
T PS::Vector2<T>::operator ^ (const PS::Vector2<T> & rhs) const;
```

#### • Argument

rhs: Input. Type const PS::Vector2<T> &.

#### • Return value

Type T. Take outer product of it and rhs, and return the results.

#### 6.4.2.9 Typecast to Vector2<U>

```
template<typename T>
template <typename U>
PS::Vector2<T>::operator PS::Vector2<U> () const;
```

### • Argument

None.

#### • Return value

Type const PS::Vector2<U>.

#### • Feature

Typecast from type const PS::Vector2<T> to type const PS::Vector2<U>.

#### 6.4.3 PS::Vector3

PS::Vector3 has three components: x, y, and z. We define various APIs and operators for these components. In the following, we list them.

#### Listing 8: Vector3

```
1 namespace ParticleSimulator{
2
       template <typename T>
3
       class Vector3{
4
       public:
5
           // Three member variables
6
           T x, y, z;
7
8
           // Constructor
9
           Vector3() : x(T(0)), y(T(0)), z(T(0)) {}
10
           Vector3(const T _x, const T _y, const T _z) : x(_x), y(
                 _y), z(_z) {}
11
           Vector3(const T s) : x(s), y(s), z(s) {}
12
           Vector3(const Vector3 & src) : x(src.x), y(src.y), z(
                 src.z) {}
13
14
           // Assignment operator
15
           const Vector3 & operator = (const Vector3 & rhs);
16
17
           //[] operators
           const T & opertor[](const int i);
18
19
           T & operator[](const int i);
20
           // Addition and subtraction
21
22
           Vector3 operator + (const Vector3 & rhs) const;
```

```
23
           const Vector3 & operator += (const Vector3 & rhs);
24
           Vector3 operator - (const Vector3 & rhs) const;
25
           const Vector3 & operator -= (const Vector3 & rhs);
26
27
           // Scalar multiplication
28
           Vector3 operator * (const T s) const;
29
           const Vector3 & operator *= (const T s);
           friend Vector3 operator * (const T s, const Vector3 & v
30
           Vector3 operator / (const T s) const;
31
32
           const Vector3 & operator /= (const T s);
33
34
           // Inner product (scalar product)
35
           T operator * (const Vector3 & rhs) const;
36
37
           // Outer product (vector product).
38
           Vector3 operator ^ (const Vector3 & rhs) const;
39
40
           // Typecast to Vector3<U>
           template <typename U>
41
42
           operator Vector3<U> () const;
43
       };
44 }
```

# 6.4.3.1 Constructor

```
template<typename T>
PS::Vector3<T>::Vector3()
```

#### • Argument

None.

#### • Feature

Default constructor. The member variables x, y and z are initialized as 0.

```
template<typename T>
PS::Vector3<T>::Vector3(const T _x, const T _y, const T _z)
```

#### • Argument

```
_x: Input. Type const T._y: Input. Type const T._z: Input. Type const T.
```

#### • Feature

Values of members x, y and z are set to values of arguments \_x, \_y and \_z, respectively.

```
template<typename T>
PS::Vector3<T>::Vector3(const T s);
```

# • Argument

s: Input. Type const T.

#### • Feature

Values of members x, y and z are set to the value of argument s.

# 6.4.3.2 Copy constructor

```
template<typename T>
PS::Vector3<T>::Vector3(const PS::Vector3<T> & src)
```

#### • Argument

src: Input. Type const PS::Vector3<T> &.

#### • Feature

Copy constructor. The new variable will have the same value as src.

#### 6.4.3.3 Member variables

```
template<typename T>
T PS::Vector3<T>::x;

template<typename T>
T PS::Vector3<T>::y;

template<typename T>
T PS::Vector3<T>::z;
```

#### • function

Member variables, x, y and z can be directly handled.

### 6.4.3.4 Assignment operator

### • Argument

rhs: Input. Type const PS::Vector3<T> &.

• Return value Type const PS::Vector3<T> &. Assigns the values of components of rhs to its components, and returns the vector itself.

# **6.4.3.5** [] operators

#### • Argument

i: input. Type const int.

#### • Return value

Type const  $\langle T \rangle$  &. It returns ith-component of the vector.

#### • Remark

This operator would be slower than the direct access of the member variables.

#### • Argument

i: input. Type const int.

#### • Return value

Type  $\langle T \rangle$  &. It returns ith-component of the vector.

#### • Remark

This operator would be slower than the direct access of the member variables.

#### 6.4.3.6 Addition and subtraction

#### • Argument

rhs: Input. Type const PS::Vector3<T> &.

#### • Return value

Type PS::Vector3<T>. Add the components of rhs and its components, and return the results.

### • Argument

rhs: Input. Type const PS::Vector3<T> &.

#### • Return value

Type const PS::Vector3<T> &. Add the components of rhs to its components, and return itself (lhs is changed).

# • 引数

rhs: Input. Type const PS::Vector3<T> &.

#### • Return value

Type PS::Vector3<T>. Subtract the components of rhs from its components, and return the results.

## • Argument

rhs: Input. Type const PS::Vector3<T> &.

#### • Return value

Type const PS::Vector3<T> &. Subtract the components of rhs from its components, and return itself (lhs is changed).

## 6.4.3.7 Scalar multiplication

```
template<typename T>
PS::Vector3<T> PS::Vector3<T>::operator * (const T s) const;
```

### • Argument

s: Input. Type const T.

#### • Return value

Type PS::Vector3<T>. Multiply its components by s, and return the results.

```
template<typename T>
const PS::Vector3<T> & PS::Vector3<T>::operator *= (const T s);
```

## • Argument

rhs: Input. Type const T.

## • Return value

Type const PS::Vector3<T> &. Multiply its components by s, and return itself (lhs is changed).

```
template<typename T>
PS::Vector3<T> PS::Vector3<T>::operator / (const T s) const;
```

## • Argument

s: Input. Type const T.

### • Return value

Type PS::Vector3<T>. Divide its components by s, and return the results.

```
template<typename T>
const PS::Vector3<T> & PS::Vector3<T>::operator /= (const T s);
```

## • Argument

rhs: Input. Type const T.

#### • Return value

Type const PS::Vector3 < T > &. Divide its components by s, and return itself (lhs is changed).

## 6.4.3.8 Inner and outer product

```
template<typename T>
T PS::Vector3<T>::operator * (const PS::Vector3<T> & rhs) const;
```

### • Argument

rhs: Input. Type const PS::Vector3<T> &.

#### • Return value

Type T. Take inner product of it and rhs, and return the result.

```
template<typename T>
T PS::Vector3<T>::operator ^ (const PS::Vector3<T> & rhs) const;
```

## • Argument

rhs: Input. Type const PS::Vector3<T> &.

#### • Return value

Type T. Take outer product of it and rhs, and return the results.

## 6.4.3.9 Typecast to Vector3<U>

```
template<typename T>
template <typename U>
PS::Vector3<T>::operator PS::Vector3<U> () const;
```

## • Argument

None.

## • Return value

Type const PS::Vector3<U>.

#### • Feature

Typecast from type const PS::Vector3<T> to const PS::Vector3<U>.

## 6.4.4 Wrappers

The wrappers of vector types are defined as follows.

Listing 9: vectorwrapper

```
namespace ParticleSimulator{
1
2
       typedef Vector2<F32> F32vec2;
3
       typedef Vector3<F32> F32vec3;
4
       typedef Vector2<F64> F64vec2;
5
       typedef Vector3<F64> F64vec3;
6
  #ifdef PARTICLE_SIMULATOR_TWO_DIMENSION
7
       typedef F32vec2 F32vec;
8
       typedef F64vec2 F64vec;
9
  #else
10
       typedef F32vec3 F32vec;
11
       typedef F64vec3 F64vec;
12 #endif
13 }
```

PS::F32vec2, PS::F32vec3, PS::F64vec2, and PS::F64vec3 are, respectively, 2D vector in single precision, 3D vector in single presicion, 2D vector in double precision, and 3D vector in double precision. If users set 2D (3D) coordinate system, PS::F32vec and PS::F64vec is wrappers of PS::F32vec2 and PS::F64vec2 (PS::F32vec3 and PS::F64vec3).

# 6.5 Orthotope type

#### 6.5.1 Abstract

The orthotope types are PS::Orthotope2 (rectangle) and PS::Orthotope3 (cuboid). We describe these orthotope types first. These are template classes, which can take basic datatypes as F32 or F64 as template arguments. We then present wrappers for these orthotope types.

## 6.5.2 PS::Orthotope2

PS::Orthotope2 has two components: low\_ and high\_, both of which are PS::Vector2 class. We define various APIs and operators for these components. In the following, we describe them.

### Listing 10: Orthotope2

```
namespace ParticleSimulator{
1
2
      template < class T>
3
      class Orthotope2{
4
      public:
           Vector2 < T > low_;
5
6
           Vector2<T> high_;
7
8
           Orthotope2(): low_(9999.9), high_(-9999.9){}
9
```

```
Orthotope2(const Vector2<T> & _low, const Vector2<T> &
10
                 _high)
               : low_(_low), high_(_high){}
11
12
           Orthotope2(const Orthotope2 & src) : low_(src.low_),
13
                 high_(src.high_){}
14
           Orthotope2(const Vector2<T> & center, const T length) :
15
               low_(center-(Vector2<T>)(length)), high_(center+(
16
                     Vector2<T>)(length)) {
           }
17
18
19
           void initNegativeVolume(){
20
               low_ = std::numeric_limits<float>::max() / 128;
21
               high_{-} = -low_{-};
           }
22
23
           void init(){
24
25
               initNegativeVolume();
26
           }
27
28
           void merge( const Orthotope2 & ort ){
29
               this->high_.x = ( this->high_.x > ort.high_.x ) ?
                     this->high_.x : ort.high_.x;
30
               this->high_.y = ( this->high_.y > ort.high_.y ) ?
                     this->high_.y : ort.high_.y;
               this->low_.x = (this->low_.x <= ort.low_.x)?
31
                     this->low_.x : ort.low_.x;
32
               this->low_.y = ( this->low_.y <= ort.low_.y ) ?
                     this->low_.y : ort.low_.y;
33
           }
34
           void merge( const Vector2<T> & vec ){
35
               this->high_.x = ( this->high_.x > vec.x ) ? this->
36
                     high_.x : vec.x;
37
               this->high_.y = ( this->high_.y > vec.y ) ? this->
                     high_.y : vec.y;
38
               this->low_.x = ( this->low_.x <= vec.x ) ? this->
                     low_.x : vec.x;
39
               this->low_.y = ( this->low_.y <= vec.y ) ? this->
                     low_.y : vec.y;
           }
40
41
           void merge( const Vector2<T> & vec, const T size){
42
43
               this \rightarrow high_.x = (this \rightarrow high_.x > vec.x + size)?
                     this->high_.x : vec.x + size;
```

### 6.5.2.1 Member variables

```
template<typename T>
PS::Vector2<T> PS::Orthotope2<T>::low_;

template<typename T>
PS::Vector2<T> PS::Orthotope2<T>::high_;
```

#### • Feature

Member variables, low\_ and high\_ can be directly handled.

### 6.5.2.2 Constructors

```
template<typename T>
PS::Orthotope2<T>::Orthotope2();
```

#### • Arguments

None. But, the template argument T must be either PS::F32 or PS::F64.

#### • Feature

Default constructor. Member variables low\_ and high\_ are initialized by (9999.9, 9999.9) and (-9999.9, -9999.9), respectively.

```
template<typename T>
PS::Orthotope2<T>::Orthotope2(const Vector2<T> _low, const Vector2<T> _high);
```

## • Arguments

```
_low: Input. Type const Vector2<T>.
_high: Input. Type const Vector2<T>.
where T must be either PS::F32 or PS::F64.
```

#### • Feature

Member variables low\_ and high\_ are initialized by \_low and \_high, respectively.

```
template<typename T>
PS::Orthotope2<T>::Orthotope2(const Vector2<T> & center, const T length);
```

### • Arguments

center: Input. Type const Vector2<T> &. length: Input. Type const T.

#### • Feature

Member variables low\_ and high\_ are initialized by center-(Vector2<T>) (length) and center+(Vector2<T>) (length), respectively.

## 6.5.2.3 Copy constructor

```
template<typename T>
PS::Orthotope2<T>::Orthotope2(const Orthotope2<T> & src);
```

## • Argument

src: Input. Type const Orthotope2<T> &.

#### • Feature

Member variables low\_ and high\_ are initialized by src.low\_ and src.high\_, respectively.

#### 6.5.2.4 Initialize

```
template<typename T>
PS::Orthotope2<T>::initNegativeVolume();
```

## • Arguments

None.

#### • Feature

Member variables  $low_a$  and  $high_a$  are initialized by (a, a) and (-a, -a), respectively, where  $a=std::numeric_limits < float>::max() / 128.$ 

```
template<typename T>
PS::Orthotope2<T>::init();
```

## • Arguments

None.

#### • Feature

This member function is the same as initNegativeVolume.

## 6.5.2.5 Merge operations

```
template<typename T>
void PS::Orthotope2<T>::merge( const Orthotope2 & ort )();
```

## • Arguments

ort: Input. Type const Orthotope2 &.

#### • Feature

Member variables low\_ and high\_ are updated so that they represents the smallest rectangle that contains both the original rectangle and rectangle ort.

```
template<typename T>
void PS::Orthotope2<T>::merge( const Vector2<T> & vec )();
```

#### • Arguments

vec: Input. Type const Vector2<T> &.

#### • Feature

Member variables low\_ and high\_ are updated so that they represents the smallest rectangle that contains the point vec.

```
template<typename T>
void PS::Orthotope2<T>::merge( const Vector2<T> & vec, const T size )();
```

### • Arguments

vec: Input. Type const Vector2<T> &. size: Input. Type const T.

### • Feature

Member variables low\_ and high\_ are updated so that they represents the smallest rectangle that contains a circle whose center is vec and radius is size.

### 6.5.3 PS::Orthotope3

PS::Orthotope3 has two components: low\_ and high\_, both of which are PS::Vector3 class. We define various APIs and operators for these components. In the following, we describe them.

Listing 11: Orthotope3

```
1 namespace ParticleSimulator{
2
       template < class T>
3
       class Orthotope3{
4
       public:
5
           Vector3<T> low_;
6
           Vector3<T> high_;
7
8
           Orthotope3(): low_(9999.9), high_(-9999.9){}
9
           Orthotope3(const Vector3<T> & _low, const Vector3<T> &
10
                 _high)
11
               : low_(_low), high_(_high) {}
12
           Orthotope3(const Orthotope3 & src) : low_(src.low_),
13
                 high_(src.high_){}
14
           Orthotope3(const Vector3<T> & center, const T length) :
15
               low_(center-(Vector3<T>)(length)), high_(center+(
16
                     Vector3<T>)(length)) {
           }
17
18
19
           void initNegativeVolume(){
               low_ = std::numeric_limits<float>::max() / 128;
20
21
               high_{-} = -low_{-};
22
           }
23
           void init(){
24
25
               initNegativeVolume();
26
           }
27
28
           void merge( const Orthotope3 & ort ){
29
               this->high_.x = ( this->high_.x > ort.high_.x ) ?
                     this->high_.x : ort.high_.x;
30
               this->high_.y = ( this->high_.y > ort.high_.y ) ?
                     this->high_.y : ort.high_.y;
31
               this->high_.z = ( this->high_.z > ort.high_.z ) ?
                     this->high_.z : ort.high_.z;
32
               this->low_.x = (this->low_.x <= ort.low_.x)?
                     this->low_.x : ort.low_.x;
33
               this->low_.y = ( this->low_.y <= ort.low_.y ) ?
```

```
this->low_.y : ort.low_.y;
                this->low_.z = ( this->low_.z <= ort.low_.z ) ?
34
                      this->low_.z : ort.low_.z;
35
            }
36
37
            void merge( const Vector3<T> & vec ){
                this \rightarrow high_.x = (this \rightarrow high_.x > vec.x)? this \rightarrow
38
                      high_.x : vec.x;
                this->high_.y = ( this->high_.y > vec.y ) ? this->
39
                      high_.y : vec.y;
40
                this->high_.z = ( this->high_.z > vec.z ) ? this->
                      high_.z : vec.z;
41
                this->low_.x = ( this->low_.x <= vec.x ) ? this->
                      low_.x : vec.x;
                this->low_.y = ( this->low_.y <= vec.y ) ? this->
42
                      low_.y : vec.y;
43
                this->low_.z = ( this->low_.z <= vec.z ) ? this->
                      low_.z : vec.z;
44
            }
45
            void merge( const Vector3<T> & vec, const T size){
46
                this \rightarrow high_.x = (this \rightarrow high_.x > vec.x + size)?
47
                      this->high_.x : vec.x + size;
48
                this \rightarrow high_.y = (this \rightarrow high_.y > vec.y + size)?
                      this->high_.y : vec.y + size;
49
                this \rightarrow high_.z = (this \rightarrow high_.z > vec.z + size)?
                      this->high_.z : vec.z + size;
                this->low_.x = ( this->low_.x <= vec.x - size) ?
50
                      this->low_.x : vec.x - size;
51
                this->low_.y = ( this->low_.y <= vec.y - size) ?
                      this->low_.y : vec.y - size;
                this->low_.z = ( this->low_.z <= vec.z - size) ?
52
                      this->low_.z : vec.z - size;
53
            }
54
       };
55 }
56 namespace PS = ParticleSimulator;
```

#### 6.5.3.1 Member variables

```
template<typename T>
PS::Vector3<T> PS::Orthotope3<T>::low_;

template<typename T>
PS::Vector3<T> PS::Orthotope3<T>::high_;
```

#### • Feature

Member variables, low\_ and high\_ can be directly handled.

#### 6.5.3.2 Constructors

```
template<typename T>
PS::Orthotope3<T>::Orthotope3();
```

## • Arguments

None. But, the template argument T must be either PS::F32 or PS::F64.

#### • Feature

Default constructor. Member variables low\_ and high\_ are initialized by (9999.9, 9999.9, 9999.9) and (-9999.9, -9999.9, -9999.9), respectively.

```
template<typename T>
PS::Orthotope3<T>::Orthotope3(const Vector3<T> _low, const Vector3<T> _high);
```

## • Arguments

```
_low: Input. Type const Vector3<T>.
_high: Input. Type const Vector3<T>.
where T must be either PS::F32 or PS::F64.
```

#### • Feature

Member variables low\_ and high\_ are initialized by \_low and \_high, respectively.

```
template<typename T>
PS::Orthotope3<T>::Orthotope3(const Vector3<T> & center, const T length);
```

## • Arguments

center: Input. Type const Vector3<T> &. length: Input. Type const T.

#### • Feature

Member variables low\_ and high\_ are initialized by center-(Vector3<T>) (length) and center+(Vector3<T>) (length), respectively.

## 6.5.3.3 Copy constructor

```
template<typename T>
PS::Orthotope3<T>::Orthotope3(const Orthotope3<T> & src);
```

## • Argument

src: Input. Type const Orthotope3<T> &.

#### • Feature

Member variables low\_ and high\_ are initialized by src.low\_ and src.high\_, respectively.

#### 6.5.3.4 Initialize

```
template<typename T>
PS::Orthotope3<T>::initNegativeVolume();
```

## • Arguments

None.

## • Feature

Member variables  $low_a$  and  $high_a$  are initialized by (a, a, a) and (-a, -a, -a), respectively, where  $a=std::numeric_limits < float>::max() / 128.$ 

```
template<typename T>
PS::Orthotope3<T>::init();
```

## • Arguments

None.

## • Feature

This member function is the same as initNegativeVolume.

## 6.5.3.5 Merge operations

```
template<typename T>
void PS::Orthotope3<T>::merge( const Orthotope3 & ort )();
```

## • Arguments

ort: Input. Type const Orthotope3 &.

#### • Feature

Member variables low\_ and high\_ are updated so that they represents the smallest cuboid that contains both the original cuboid and cuboid ort.

```
template<typename T>
void PS::Orthotope3<T>::merge( const Vector3<T> & vec )();
```

### • Arguments

vec: Input. Type const Vector3<T> &.

#### • Feature

Member variables low\_ and high\_ are updated so that they represents the smallest cuboid that contains the point vec.

```
template<typename T>
void PS::Orthotope3<T>::merge( const Vector3<T> & vec, const T size )();
```

### • Arguments

vec: Input. Type const Vector3<T> &. size: Input. Type const T.

#### • Feature

Member variables low\_ and high\_ are updated so that they represents the smallest cuboid that contains a sphere whose center is vec and radius is size.

## 6.5.4 Wrappers

The wrappers of orthotope types are defined as follows.

## Listing 12: Orthotope wrapper

```
1 namespace ParticleSimulator{
2
       typedef Orthotope2<F32> F32ort2;
3
       typedef Orthotope3<F32> F32ort3;
       typedef Orthotope2<F64> F64ort2;
4
5
       typedef Orthotope3<F64> F64ort3;
  #ifdef PARTICLE_SIMULATOR_TWO_DIMENSION
6
7
       typedef F32ort2 F32ort;
8
       typedef F64ort2 F64ort;
9 #else
10
       typedef F32ort3 F32ort;
11
       typedef F64ort3 F64ort;
12 #endif
13 }
```

PS::F32ort2, PS::F32ort3, PS::F64ort2, and PS::F64ort3 are, respectively, 2D orthotope in single precision, 3D orthotope in single presicion, 2D orthotope in double precision, and 3D orthotope in double precision. If users set 2D (3D) coordinate system, PS::F32ort and PS::F64ort are wrappers of PS::F32ort2 and PS::F64ort2 (PS::F32ort3 and PS::F64ort3).

# 6.6 Symmetric matrix type

#### 6.6.1 Abstract

Symmetric matrix types are PS::MatrixSym2 (2x2 matrix) and PS::MatrixSym3 (3x3 matrix). We describe these matrix types first and then, we present wrappers for these matrix types.

### 6.6.2 PS::MatrixSym2

PS::MatrixSym2 has three components: xx, yy, and xy. We define various APIs and operators for these components. In the following, we list them.

### Listing 13: MatrixSym2

```
1 namespace ParticleSimulator{
2
       template < class T>
3
       class MatrixSym2{
4
       public:
5
           // Three member variables
6
           T xx, yy, xy;
7
8
           // Constructors
           MatrixSym2() : xx(T(0)), yy(T(0)), xy(T(0)) {}
9
           MatrixSym2(const T _xx, const T _yy, const T _xy)
10
               : xx(_xx), yy(_yy), xy(_xy) {}
11
           MatrixSym2(const T s) : xx(s), yy(s), xy(s){}
12
13
           MatrixSym2(const MatrixSym2 & src) : xx(src.xx), yy(src
                 .yy), xy(src.xy) {}
14
15
           // Assignment operator
           const MatrixSym2 & operator = (const MatrixSym2 & rhs);
16
17
           // Addition and subtraction
18
19
           MatrixSym2 operator + (const MatrixSym2 & rhs) const;
           const MatrixSym2 & operator += (const MatrixSym2 & rhs)
20
           MatrixSym2 operator - (const MatrixSym2 & rhs) const;
21
22
           const MatrixSym2 & operator -= (const MatrixSym2 & rhs)
                  const;
23
24
           // Trace
25
           T getTrace() const;
```

#### 6.6.2.1 Constructor

```
template<typename T>
PS::MatrixSym2<T>::MatrixSym2();
```

## • Argument

None.

#### • Feature

Default constructor. The member variables xx, yy and xy are initialized as 0.

## • Argument

```
_xx: Input. Type const T.
_yy: Input. Type const T.
_xy: Input. Type const T.
```

#### • Feature

Values of members xx, yy and xy are set to values of arguments  $_xx$ ,  $_yy$  and  $_xy$ , respectively.

```
template<typename T>
PS::MatrixSym2<T>::MatrixSym2(const T s);
```

## • Argument

s: Input. Type const T.

#### • Feature

Values of members xx, yy and xy are set to the value of argument s.

## 6.6.2.2 Copy constructor

```
template<typename T>
PS::MatrixSym2<T>::MatrixSym2(const PS::MatrixSym2<T> & src)
```

### • Argument

```
src: Input. Type const PS::MatrixSym2<T> &.
```

#### • Feature

Copy constructor. The new variable will have the same value as src.

## 6.6.2.3 Assignment operator

## • Argument

rhs: Input. Type const PS::MatrixSym2<T> &.

## • Return value

Type const PS::MatrixSym2<T> &. Assigns the values of components of rhs to its components, and returns the vector itself.

#### 6.6.2.4 Addition and subtraction

#### • Argument

```
rhs: Input. Type const PS::MatrixSym2<T> &.
```

#### • Return value

Type PS::MatrixSym2 < T>. Add the components of rhs and its components, and return the results.

## • Argument

rhs: Input. Type const PS::MatrixSym2<T> &.

#### • Return value

Type const PS::MatrixSym2<T> &. Add the components of rhs to its components, and return itself (lhs is changed).

## • Argument

rhs: Input. Type const PS::MatrixSym2<T> &.

#### • Return value

Type PS::MatrixSym2 < T>. Subtract the components of rhs from its components, and return the results.

## • Argument

rhs: Input. Type const PS::MatrixSym2<T> &.

#### • Return value

Type const PS::MatrixSym2<T> &. Subtract the components of rhs from its components, and return itself (lhs is changed).

#### 6.6.2.5 Trace

```
template<typename T>
T PS::MatrixSym2<T>::getTrace() const;
```

## • Argument

None.

#### • Return value

Type T.

#### • Feature

Calculate the trace, and return the result.

## 6.6.2.6 Typecast to MatrixSym2<U>

```
template<typename T>
template<typename U>
PS::MatrixSym2<T>::operator PS::MatrixSym2<U> () const;
```

### • Argument

None.

#### • Return value

Type const PS::MatrixSym2<U>.

#### • Feature

Typecast from type const PS::MatrixSym2<T> to const PS::MatrixSym2<U>.

## 6.6.3 PS::MatrixSym3

PS::MatrixSym3 has six components: xx, yy, zz, xy, xz, and yz. We define various APIs and operators for these components. In the following, we list them.

Listing 14: MatrixSym3

```
1 namespace ParticleSimulator{
2
       template < class T>
3
       class MatrixSym3{
4
       public:
5
           // Six member variables
6
           T xx, yy, zz, xy, xz, yz;
7
8
           // Constructors
           MatrixSym3() : xx(T(0)), yy(T(0)), zz(T(0)),
9
10
                           xy(T(0)), xz(T(0)), yz(T(0)) {}
           MatrixSym3(const T _xx, const T _yy, const T _zz,
11
                       const T _xy, const T _xz, const T _yz )
12
                           : xx(_xx), yy(_yy), zz(_zz),
13
14
                           xy(_xy), xz(_xz), yz(_yz) {}
           MatrixSym3(const T s) : xx(s), yy(s), zz(s),
15
16
                                    xy(s), xz(s), yz(s) {}
17
           MatrixSym3(const MatrixSym3 & src) :
18
               xx(src.xx), yy(src.yy), zz(src.zz),
               xy(src.xy), xz(src.xz), yz(src.yz) {}
19
20
21
           // Assignment operator
22
           const MatrixSym3 & operator = (const MatrixSym3 & rhs);
23
24
           // Addition and subtraction
```

```
25
           MatrixSym3 operator + (const MatrixSym3 & rhs) const;
           const MatrixSym3 & operator += (const MatrixSym3 & rhs)
26
                  const;
27
           MatrixSym3 operator - (const MatrixSym3 & rhs) const;
28
           const MatrixSym3 & operator -= (const MatrixSym3 & rhs)
                  const;
29
           // Trace
30
31
           T getTrace() const;
32
33
           // Typecast to MatrixSym3 < U>
34
           template <typename U>
35
           operator MatrixSym3<U> () const;
       }
36
37 }
38 namespace PS = ParticleSimulator;
```

### 6.6.3.1 Constructor

```
template<typename T>
PS::MatrixSym3<T>::MatrixSym3();
```

## • Argument

None.

#### • Feature

Default constructor. All the member variables are initialized as 0.

## • Argument

```
_xx: Input. Type const T.
_yy: Input. Type const T.
_zz: Input. Type const T.
_xy: Input. Type const T.
```

```
_xz: Input. Type const T.
_yz: Input. Type const T.
```

#### • Feature

Values of members xx, yy, zz, xy, xz and yz are set to values of arguments \_xx, \_yy, \_zz, \_xy, xz and \_yz, respectively.

```
template<typename T>
PS::MatrixSym3<T>::MatrixSym3(const T s);
```

## • Argument

s: Input. Type const T.

#### • Feature

Values of members are set to the value of argument s.

## 6.6.3.2 Copy constructor

```
template<typename T>
PS::MatrixSym3<T>::MatrixSym3(const PS::MatrixSym3<T> & src)
```

#### • Argument

```
src: Input. Type const PS::MatrixSym3<T> &.
```

## • Feature

Copy constructor. The new variable will have the same value as src.

## 6.6.3.3 Assignment operator

#### • Argument

```
rhs: Input. Type const PS::MatrixSym3<T> &.
```

## • Return value

Type const PS::MatrixSym3<T> &. Assigns the values of components of rhs to its components, and returns the vector itself.

## 6.6.3.4 Addition and subtraction

### • Argument

rhs: Input. Type const PS::MatrixSym3<T> &.

### • Return value

Type PS::MatrixSym3 < T>. Add the components of rhs and its components, and return the results.

### • Argument

rhs: Input. Type const PS::MatrixSym3<T> &.

#### • Return value

Type const PS::MatrixSym3<T> &. Add the components of rhs to its components, and return itself (lhs is changed).

## • Argument

rhs: Input. Type const PS::MatrixSym3<T> &.

#### • Return value

Type PS::MatrixSym3 < T>. Subtract the components of rhs from its components, and return the results.

### • Argument

rhs: Input. Type const PS::MatrixSym3<T> &.

## • Return value

Type const PS::MatrixSym3<T> &. Subtract the components of rhs from its components, and return itself (lhs is changed).

#### 6.6.3.5 Trace

```
template<typename T>
T PS::MatrixSym3<T>::getTrace() const;
```

# • Argument

None.

#### • Return value

Type T.

#### • Feature

Calculate the trace, and return the result.

## 6.6.3.6 Typecast to MatrixSym3<U>

```
template<typename T>
template<typename U>
PS::MatrixSym3<T>::operator PS::MatrixSym3<U> () const;
```

## • Argument

None.

#### • Return value

Type const PS::MatrixSym3<U>.

#### • Feature

Typecast from type const PS::MatrixSym3<T> to const PS::MatrixSym3<U>.

## 6.6.4 Wrappers

The wrappers of symmetric matrix types are defined as follows.

## Listing 15: matrixsymwrapper

```
1 namespace ParticleSimulator{
2    typedef MatrixSym2 < F32 > F32mat2;
3    typedef MatrixSym3 < F32 > F32mat3;
4    typedef MatrixSym2 < F64 > F64mat2;
5    typedef MatrixSym3 < F64 > F64mat3;
```

```
6 #ifdef PARTICLE_SIMULATOR_TWO_DIMENSION
7          typedef F32mat2 F32mat;
8          typedef F64mat2 F64mat;
9 #else
10          typedef F32mat3 F32mat;
11          typedef F64mat3 F64mat;
12 #endif
13 }
14 namespace PS = ParticleSimulator;
```

PS::F32mat2, PS::F32mat3, PS::F64mat2, and PS::F64mat3 are, respectively, 2x2 symmetric matrix in single precision, 3x3 symmetric matrix in single precision, 2x2 symmetric matrix in double precision, and 3x3 symmetric matrix in double precision. If users set 2D (3D) coordinate system, PS::F32mat and PS::F64mat is wrappers of PS::F32mat2 and PS::F64mat2 (PS::F32mat3 and PS::F64mat3).

# 6.7 PS::SEARCH\_MODE type

## 6.7.1 Summary

In this section, we describe data type PS::SEARCH\_MODE. It is used only as template arguments of class PS::TreeForForce. This data type determines the interaction mode of the class. PS::SEARCH\_MODE can take the following values:

- PS::SEARCH\_MODE\_LONG
- PS::SEARCH\_MODE\_LONG\_CUTOFF
- PS::SEARCH\_MODE\_GATHER
- PS::SEARCH\_MODE\_SCATTER
- PS::SEARCH\_MODE\_SYMMETRY
- PS::SEARCH\_MODE\_LONG\_SCATTER
- PS::SEARCH\_MODE\_LONG\_SYMMETRY
- PS::SEARCH\_MODE\_LONG\_CUTOFF\_SCATTER

Each of them corresponds to a mode for interaction calculation. In the following, we describe them.

#### 6.7.2 PS::SEARCH\_MODE\_LONG

This type is used when a group of distant particles is regarded as a superparticle as in the standard Barnes-Hut tree code. This type is for gravitational force and Coulomb's force under open boundary condition.

#### 6.7.3 PS::SEARCH\_MODE\_LONG\_CUTOFF

This type is used when a group of distant particles is regarded as a superparticle, and when its force does not reach to infinity. This type is for gravitational force and Coulomb's force under periodic boundary condition.

#### 6.7.4 PS::SEARCH\_MODE\_GATHER

This type is used when its force decays to zero at a finite distance, and when the distance is determined by the size of *i*-particle.

## 6.7.5 PS::SEARCH\_MODE\_SCATTER

This type is used when its force decays to zero at a finite distance, and when the distance is determined by the size of j-particle.

#### 6.7.6 PS::SEARCH\_MODE\_SYMMETRY

This type is used when its force decays to zero at a finite distance, and when the distance is determined by the larger of the sizes of i- and j-particles.

### 6.7.7 PS::SEARCH\_MODE\_LONG\_SCATTER

Almost the same as SEARCH\_MODE\_LONG, but if the distance between i- and j-particles is smaller than the search radius of j-particle, the j-particle is not included in superparticle.

#### 6.7.8 PS::SEARCH\_MODE\_LONG\_SYMMETRY

Almost the same as SEARCH\_MODE\_LONG, but if the distance between i- and j-particles is smaller than the larger of the search radii of i- and j-particle, the j-particle is not included in superparticle.

## 6.7.9 PS::SEARCH\_MODE\_LONG\_CUTOFF\_SCATTER

Not implemented yet.

# 6.8 Enumerated type

## 6.8.1 Summary

In this section, we describe enumerated types defined in FDPS. Currently, there is just one datatype. We describe it below.

## 6.8.2 PS::BOUNDARY\_CONDITION type

#### 6.8.2.1 Summary

Type BOUNDARY\_CONDITION specifies boundary conditions. The definition is as follows.

## Listing 16: boundarycondition

1 namespace ParticleSimulator{
2 enum BOUNDARY\_CONDITION{
3 BOUNDARY\_CONDITION\_OPEN,
4 BOUNDARY\_CONDITION\_PERIODIC\_X,

```
5
           BOUNDARY_CONDITION_PERIODIC_Y,
6
           BOUNDARY_CONDITION_PERIODIC_Z,
7
           BOUNDARY_CONDITION_PERIODIC_XY,
8
           BOUNDARY_CONDITION_PERIODIC_XZ,
9
           BOUNDARY_CONDITION_PERIODIC_YZ,
10
           BOUNDARY_CONDITION_PERIODIC_XYZ,
           BOUNDARY_CONDITION_SHEARING_BOX,
11
12
           BOUNDARY_CONDITION_USER_DEFINED,
13
       };
14 }
```

We explain each value below.

#### 6.8.2.2 PS::BOUNDARY\_CONDITION\_OPEN

This specifies the open boundary condition.

#### 6.8.2.3 PS::BOUNDARY\_CONDITION\_PERIODIC\_X

This specifies the periodic boundary condition in the direction of x-axis, and open boundary condition in other directions. The interval is left-bounded and right-unbounded. This is true for all periodic boundary conditions.

#### 6.8.2.4 PS::BOUNDARY\_CONDITION\_PERIODIC\_Y

This specifies the periodic boundary condition in the direction of y-axis, and open boundary condition in other directions.

#### 6.8.2.5 PS::BOUNDARY\_CONDITION\_PERIODIC\_Z

This specifies the periodic boundary condition in the direction of z-axis, and open boundary condition in other directions.

#### 6.8.2.6 PS::BOUNDARY\_CONDITION\_PERIODIC\_XY

This specifies the periodic boundary condition in the directions of x- and y-axes, and open boundary condition in the direction of z-axis.

#### 6.8.2.7 PS::BOUNDARY\_CONDITION\_PERIODIC\_XZ

This specifies the periodic boundary condition in the directions of x- and z-axes, and open boundary condition in the direction of y-axis.

## 6.8.2.8 PS::BOUNDARY\_CONDITION\_PERIODIC\_YZ

This specifies the periodic boundary condition in the directions of y- and z-axes, and open boundary condition in the direction of x-axis.

#### 6.8.2.9 PS::BOUNDARY\_CONDITION\_PERIODIC\_XYZ

This specifies the periodic boundary condition in all three directions.

#### 6.8.2.10 PS::BOUNDARY\_CONDITION\_SHEARING\_BOX

Not implemented yet.

#### 6.8.2.11 PS::BOUNDARY\_CONDITION\_USER\_DEFINED

Not implemented yet.

### 6.8.3 PS::INTERACTION\_LIST\_MODE type

## 6.8.3.1 Summary

Type INTERACTION\_LIST\_MODE is used to determine if user program reuse the interaction list or not. This type is defined as follows.

## Listing 17: boundarycondition

```
namespace ParticleSimulator{
    enum INTERACTION_LIST_MODE{
        MAKE_LIST,
        MAKE_LIST_FOR_REUSE,
        REUSE_LIST,
    };
}
```

This data type is used as the last argument of the function calcForceAllAndWriteBack(). For more detail, plese see section ??.

## 6.8.3.2 PS::MAKE\_LIST

FDPS (re)makes interaction lists for each interaction calculation (each call of the APIs described above). In this case, we cannot reuse interaction list in the next interaction calculation because FDPS does not store the information of interaction list. **This is the default operation mode in FDPS**.

# 6.8.3.3 PS::MAKE\_LIST\_FOR\_REUSE

FDPS (re)makes interaction lists and stores them internally. Then, it performs interaction calculation. In this case, we can reuse these interaction lists in the next interaction calculation if we call the APIs with the flag PS::REUSE\_LIST. The interaction lists

memorized in FDPS are destroyed if we perform the interaction calculation with the flags PS::MAKE\_LIST\_FOR\_REUSE or PS::MAKE\_LIST.

#### 6.8.3.4 PS::REUSE\_INTERACTION\_LIST

FDPS performs interaction calculation using the previously-created interaction lists, which are the lists that are created at the previous call of the APIs with the flag PS::MAKE\_LIST\_FOR\_REU In this case, moment information in superparticles are automatically updated using the latest particle information.

# 6.9 PS::TimeProfile type

#### 6.9.1 Abstract

In this section, we describe data type PS::TimeProfile. This data type is class to store calculation time for each function, and is used for three classes: DomainInfo, ParticleSystem, and TreeForForce. These three classes have "PS::TimeProfile getTimeProfile()". Users get the calculation time of each function by using the function "getTimeProfile()".

This class is described as follows.

Listing 18: TimeProfile

```
namespace ParticleSimulator{
1
2
       class TimeProfile{
3
       publid:
4
           F64 collect_sample_particle;
5
           F64 decompose_domain;
6
           F64 exchange_particle;
7
           F64 make_local_tree;
8
           F64 make_global_tree;
9
           F64 calc_force;
10
           F64 calc_moment_local_tree;
11
           F64 calc_moment_global_tree;
12
           F64 make_LET_1st;
13
           F64 make_LET_2nd;
14
           F64 exchange_LET_1st;
15
           F64 exchange_LET_2nd;
16
       };
17 }
```

#### 6.9.1.1 Addition

```
PS::TimeProfile PS::TimeProfile::operator + (const PS::TimeProfile & rhs) const;
```

## • Argument

rhs: Input. Type const TimeProfile &.

#### • Return value

Type PS::TimeProfile. Add the components of rhs to its own components, and return the results.

## 6.9.1.2 Reduction

```
PS::F64 PS::TimeProfile::getTotalTime() const;
```

## • Argument

None.

## • Return value

Type PS::F64. Return values of all the member variables.

## 6.9.1.3 Initialize

```
void PS::TimeProfile::clear();
```

## • Argument

None.

#### • Return value

None.

## • Feature

Assign 0 to all the member variables.

# 7 Classes and Functors User-Defined

# 7.1 Summary

In this section we provide details of user-defined classes and functors. The user-defined classes are FullParticle, EssentialParticleI, EssentialParticleJ, Moment, SuperParticleJ, Force and Header classes. The functors are calcForceEpEp and calcForceSpEp.

A FullParticle class contains all informations of a particle and is a template argument of FDPS-defined classes (see step 0 in Sec. 2.3).

Functors calcForceEpEp and calcForceSpEp define the interaction between two EssentialParticle classes and SuperParticles class and EssentialParticle, respectively. These functors are argument of FDPS-defined TreeForForce (see step 0 in Sec. 2.3). If neither PS::SEARCH\_MODE\_LONG nor PS::SEARCH\_MODE\_LONG\_CUTOFF are used, users do not need to define calcForceSpEp.

Classes EssentialParticleI, EssentialParticleJ, Moment, SuperParticleJ and Force are support classes for the interactions between two particles. Classes EssentialParticleI and EssentialParticleJ contain the quantities of i- and j- particles used for the calculation of interactions. A Force class contains the quantities of an i- particle used to store the results of the calculations of interactions. Since these classes contain part of information of FullParticle, it is possible to use FullParticle in place of these classes. However, FullParticle may contain other values which are not used to evaluate the calculations of interactions. It is recommended to use these classes when high performance is desirable. Classes Moment and SuperParticleJ contain the information of moment and super particle, respectively. These classes are necessary if users require the super particle, i.e., when PS::SEARCH\_MODE\_LONG or PS::SEARCH\_MODE\_LONG\_CUTOFF are used. A class Header contains the header informations of input/output file.

The rest of this chapter describes how these classes and functors should be implemented. Users must write data transfer functions between these classes and interaction calculation in functor, which are used in the above member functions and functors.

## 7.2 FullParticle

# 7.2.1 Summary

The FullParticle class contains all information of a particle and is one of the template parameters of FDPS-defined ParticleSystem (see step 0 in Sec. 2.3). Users can define arbitrary member variables and member functions, as far as required member functions are defined. Below, we describe the required member functions.

## 7.2.2 Premise

Let us take FP class as an example of FullParticle as below. Users can use an arbitrary name in place of FP.

class FP;

## 7.2.3 Required member functions

## 7.2.3.1 Summary

The member functions FP::getPos and FP::copyFromForce are required. FP::getPos returns the position of a particle. Function FP::copyFromForce copies the results of calculation back to FullParticle. The examples and descriptions for these member functions are listed below.

## 7.2.3.2 FP::getPos

```
class FP {
public:
    PS::F64vec getPos() const;
};
```

- Arguments None.
- Returns

PS::F32vec or PS::F64vec. Returns the position of FP class.

• Behaviour

Returns the member variables which contains the position of a particle.

## 7.2.3.3 FP::copyFromForce

```
class Force;

class FP {
  public:
    void copyFromForce(const Force & force);
};
```

## • Arguments

force: Input. const Force type.

• Returns

None.

## • Behaviour

Copies back the results of calculation to FP class.

## 7.2.4 Required member functions for specific cases

## 7.2.4.1 Summary

In this section we describe the member functions for specific cases listed below;

- Modes other than PS::SEARCH\_MODE\_LONG for PS::SEARCH\_MODE are used.
- APIs for file I/O in ParticleSystem are used.
- ParticleSystem::adjustPositionIntoRootDomain API is used.
- Particle Mesh class, which is an extension of FDPS, is used.

## 7.2.4.2 Modes other than PS::SEARCH\_MODE\_LONG for PS::SEARCH\_MODE are used

## 7.2.4.2.1 FP::getRSearch

```
class FP {
public:
    PS::F64 getRSearch() const;
};
```

#### • Arguments

None.

#### • Returns

PS::F32vec or PS::F64vec. Returns the value of the member variable which contains neighbor search radius in FP.

#### Behaviour

Returns the value of the member variable which contains neighbor search radius in FP.

## 7.2.4.3 APIs for file I/O in ParticleSystem are used

The member functions readAscii, writeAscii, readBinary, and writeBinary are necessary, if users use ParticleSystem::readParticleAscii, ParticleSystem::writeParticleAscii, ParticleSystem::writeParticleBinary, respectively (users can also use different names for these member functions. For more details, please see section 9.1.3.2.3). In this section we describe the rules for defining readAscii, writeAscii, readBinary, and writeBinary.

## 7.2.4.3.1 FP::readAscii

```
class FP {
public:
    void readAscii(FILE *fp);
};
```

### • Arguments

fp: FILE \* type. A file pointer of input file.

#### • Returns

None.

## • Behavior

Set the values of member variables of an instance of class FP by reading data from a ASCII file specified by file pointer fp. Data of one particle must be stored in one line. In other words, Data of one particle must end with a single new-line character  $(\n)$ .

## 7.2.4.3.2 FP::writeAscii

```
class FP {
public:
    void writeAscii(FILE *fp);
};
```

### • Arguments

fp: FILE \* type. A file pointer of output file.

### • Returns

None.

### • Behavior

Output the data of an instance of class FP to a file specified by file pointer fp in a ASCII format. Data must be written as one line. In other words, data of one particle must end with a single newline character  $(\n)$ .

## 7.2.4.3.3 *FP::readBinary*

```
class FP {
 public:
    void readBinary(FILE *fp);
};
```

## • Arguments

fp: FILE \* type. A file pointer of input file.

#### • Returns

None.

#### • Behavior

Set the values of member variables of an instance of class FP by reading data from a binary file specified by file pointer fp.

## 7.2.4.3.4 FP::writeBinary

```
class FP {
public:
    void writeBinary(FILE *fp);
};
```

## • Arguments

fp: FILE \* type. A file pointer of output file.

## • Returns

None.

## • Behavior

Output the data of an instance of class FP to a file specified by file pointer fp in a binary format.

## 7.2.4.4 ParticleSystem::adjustPositionIntoRootDomain API is used

## 7.2.4.4.1 FP::setPos

```
class FP {
  public:
    void setPos(const PS::F64vec pos_new);
};
```

#### • Arguments

pos\_new: Input. const PS::F32vec or const PS::F64vec. Modified positions of particle by FDPS.

#### • Returns

None.

### • Behaviour

Replaces the positions in FP class by those modified by FDPS.

## 7.2.4.5 Particle Mesh class, which is an extension of FDPS, is used

When Particle Mesh class is used, FP::getChargeParticleMesh and FP::copyFromForceParticleMesh be defined. Below, the rules for these functions are described.

## 7.2.4.5.1 FP::getChargeParticleMesh

```
class FP {
public:
    PS::F64 getChargeParticleMesh() const;
};
```

### • Arguments

None.

#### • Returns

PS::F32 or PS::F64. Returns the mass or the electric charge of a particle.

## 7.2.4.5.2 FP::copyFromForceParticleMesh

```
class FP {
  public:
    void copyFromForceParticleMesh(const PS::F32vec & acc_pm);
};
```

#### • Arguments

acc\_pm: const PS::F32vec or const PS::F64vec. Returns the resulting force by Particle Mesh.

#### • Returns

None.

#### • Behaviour

Writes back the resulting force by Particle Mesh to a particle.

## 7.3 EssentialParticleI

#### **7.3.1** Summary

The EssentialParticleI class should contain all information of an i- particle which is necessary to calculate interaction (see step 0 in Sec. 2.3). Class EssentialParticleI should have required member functions with specific names, as described below.

#### 7.3.2 Premise

Let us take EPI and FP classes as examples of EssentialParticleI and FullParticle as below. Users can use arbitrary names in place of EPI and FP.

```
class FP;
class EPI;
```

## 7.3.3 Required member functions

## 7.3.3.1 Summary

The member functions EPI::getPos and EPI::copyFromForce are required. EPI::getPos returns the position of a particle to FDPS. EPI::copyFromFP copies the information necessary for the interaction calculation from FullParticle. The examples and descriptions for these member functions are listed below.

## 7.3.3.2 EPI::getPos

```
class EPI {
public:
    PS::F64vec getPos() const;
};
```

## • Arguments

None.

## • Returns

PS::F64vec. Returns the position of a particle of EPI class.

## 7.3.3.3 EPI::copyFromFP

```
class FP;
class EPI {
  public:
    void copyFromFP(const FP & fp);
};
```

#### • Arguments

fp: Input. const FP & type.

#### • Returns

None.

#### • Behaviour

Copies the part of information of FP to EssnetialParticleI.

## 7.3.4 Required member functions for specific case

## 7.3.4.1 Summary

In this section we describe the member functions in the case that PS::SEARCH\_MODE\_GATHER or PS::SEARCH\_MODE\_SYMMETRY are used.

#### 7.3.4.2 Modes other than PS::SEARCH\_MODE\_LONG as PS::SEARCH\_MODE are used

### 7.3.4.2.1 EPI::getRSearch

```
class EPI {
public:
    PS::F64 getRSearch() const;
};
```

## • Arguments

None.

#### • Returns

PS::F32vec or PS::F64vec. Returns the value of the neighbour search radius in EPI.

## 7.4 EssentialParticleJ

## 7.4.1 Summary

The EssentialParticleJ class should contain all information of a j- particle which is necessary to calculate interaction (see step 0 in Sec. 2.3). This class is a subset of FullParticle (see, Sec. 7.2). Class EssentialParticleJ should have required member functions with specific names, as described below.

#### 7.4.2 Premise

Let us take EPJ and FP classes as examples of EssentialParticleJ and FullParticle as below. Users can use arbitrary names in place of EPJ and FP.

```
class FP;
class EPJ;
```

## 7.4.3 Required member functions

## 7.4.3.1 Summary

The member functions EPJ::getPos and EPJ::copyFromForce are required. EPJ::getPos returns the position of a particle to FDPS. EPJ::copyFromFP copies the information necessary for the interaction calculation from FullParticle. The examples and descriptions for these member functions are listed below.

# 7.4.3.2 EPJ::getPos

```
class EPJ {
public:
    PS::F64vec getPos() const;
};
```

## • Arguments

None.

#### • Returns

PS::F64vec. Returns the position of a particle of EPJ class.

# 7.4.3.3 EPJ::copyFromFP

```
class FP;
class EPJ {
public:
    void copyFromFP(const FP & fp);
};
```

## • Arguments

fp: Input. const FP & type.

#### • Returns

None.

#### Behaviour

Copies the part of information of FP to EPJ.

## 7.4.4 Required member functions for specific cases

## 7.4.4.1 Summary

In this section we describe the member functions in the case that the other mode than PS::SEARCH\_MODE\_LONG as PS::SEARCH\_MODE or PS::BOUNDARY\_CONDITION\_OPEN as BOUNDARY\_CONDITION

are used. For the detailed description about pre-defined Moment and SuperParticleJ classes, see corresponding section.

# 7.4.4.2 Modes other than PS::SEARCH\_MODE\_LONG as PS::SEARCH\_MODE are used

# 7.4.4.2.1 EPJ::getRSearch

```
class EPJ {
public:
    PS::F64 getRSearch() const;
};
```

# • Arguments

None.

#### • Returns

PS::F32vec or PS::F64vec. Returns the value of the member variable which contains neighbor search radius in EPJ.

# 7.4.4.3 Modes other than PS::BOUNDARY\_CONDITION\_OPEN are used BOUNDARY\_CONDITION

# 7.4.4.3.1 EPJ::setPos

```
class EPJ {
public:
    void setPos(const PS::F64vec pos_new);
};
```

# • Arguments

pos\_new: Input. const PS::F32vec or const PS::F64vec. Modified positions of particle by FDPS.

### • Returns

None.

#### • Behaviour

Replaces the positions in EPJ class by those modified by FDPS.

# 7.4.4.4 Obtian EPJ from id of particle

# 7.4.4.4.1 EPJ::getId

```
class EPJ {
public:
    PS::S64 getId();
};
```

#### arguments

void.

#### • returned value

Type PS::S64.

#### • function

This function is needed when user program uses PS::TreeForForce::getEpjFromId(). For more information, please see 9.1.4.2.7.

# 7.5 Moment

# 7.5.1 Summary

The class Moment encapsulates the values of moment and related quantities of a set of grouped particles (see, step 0 in Sec. 2.3). The examples of moment are monopole, dipole and radius of the largest particle and so on. This class is used for intermediate variables to make SuperParticleJ from EssentialParticleJ. Thus, this class has a member function that calculates the moment from EssentialParticleJ or SuperParticleJ.

Since these processes have prescribed form, FDPS has pre-defined classes. In this section we first describe the pre-defined class and then describe the rules in the case users define it.

# 7.5.2 Pre-defined class

# 7.5.2.1 Summary

FDPS has several pre-defined Moment classes, which are used if specific case are chosen as PS::SEARCH\_MODE in PS::TreeForForce. Below, we describe about the Moment class which can be chosen in each PS::SEARCH\_MODE.

#### 7.5.2.2 PS::SEARCH\_MODE\_LONG

# 7.5.2.2.1 PS::MomentMonopole

This class encapsulates the monopole moment. The reference point for the calculation of monopole is set to center of mass or center of charge.

```
namespace ParticleSimulator {
    class MomentMonopole {
    public:
        F64 mass;
        F64vec pos;
    };
}
```

- Name of the class PS::MomentMonopole
- Members and their information mass: accumulated mass or electron charge. pos: center of mass or center of charge.
- Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge and EssentialParticleJ::getPos and returns mass/electron charge and position. Users can use an arbitrary name instead of EssentialParticleJ.

# 7.5.2.2.2 PS::MomentQuadrupole

This class encapsulates the monopole and quadrupole moment. The reference point for the calculation of moments is set to the center of mass.

```
namespace ParticleSimulator {
   class MomentQuadrupole {
    public:
        F64 mass;
        F64vec pos;
        F64mat quad;
    };
}
```

- Name of the class PS::MomentQuadrupole
- Members and their information

mass: accumulated mass.

pos: center of mass.

quad: accumulated quadrupole.

• Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge and EssentialParticleJ::getPos and returns mass/electron charge and position. Users can use an arbitrary name instead of EssentialParticleJ.

### 7.5.2.2.3 PS::MomentMonopoleGeometricCenter

This class encapsulates the monopole moment. The reference point for the calculation of monopole is set to the geometric center.

- Name of the class PS::MomentMonopoleGeometricCenter
- Members and their information

```
charge: accumulated mass/charge.
```

pos: geometric center.

• Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge and EssentialParticleJ::getPos and returns mass/electron charge and position. Users can use an arbitrary name instead of EssentialParticleJ.

### 7.5.2.2.4 PS::MomentDipoleGeometricCenter

This class encapsulates the moments up to dipole. The reference point for the calculation of moments is set to geometric center.

- Name of the class PS::MomentDipoleGeometricCenter
- Members and their information

```
charge: accumulated mass/charge.
```

pos: geometric center.

dipole: dipole of the particle mass or electron charge.

#### • Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge and EssentialParticleJ::getPos and returns mass/electron charge and position. Users can use an arbitrary name instead of EssentialParticleJ.

# 7.5.2.2.5 PS::MomentQuadrupoleGeometricCenter

This class encapsulates the moments up to quadrupole. The reference point for the calculation of moments is set to geometric center.

- Name of the class PS::MomentQuadrupoleGeometricCenter
- Members and their information

```
charge: accumulated mass/charge.
```

pos: geometric center.

dipole: dipole of the particle mass or electron charge.

quadrupole: quadrupole of the particle mass or electron charge.

• Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge and EssentialParticleJ::getPos and returns mass/electron charge and position. Users can use an arbitrary name instead of EssentialParticleJ.

### 7.5.2.3 PS::SEARCH\_MODE\_LONG\_SCATTER

# 7.5.2.3.1 PS::MomentMonopoleScatter

This class encapsulates the monopole moment. The reference point for the calculation of monopole is set to center of mass or center of charge.

```
namespace ParticleSimulator {
    class MomentMonopoleScatter {
    public:
        F64 mass;
        F64vec pos;
        F64ort vertex_out_;
        F64ort vertex_in_;
    };
}
```

- Name of the class PS::MomentMonopoleScatter
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

vertex\_out\_: positions of vertices of the smallest rectangular parallelepiped including all grouped particles when each particle is regarded as a sphere whose radius is the returned value of EssentialParticleJ::getRSearch.

vertex\_in\_: positions of vertices of the smallest rectangular parallelepiped including
all grouped particles.

• Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge, EssentialParticleJ::getPos and EssentialParticleJ::getRSearch and returns mass/electron charge, position and cutoff length. Users can use an arbitrary name instead of the name of member functions of EssentialParticleJ.

# 7.5.2.3.2 PS::MomentQuadrupoleScatter

This class encapsulates the monopole and quadrupole moments. The reference point for the calculation of monopole is set to center of mass or center of charge.

```
namespace ParticleSimulator {
    class MomentQuadrupoleScatter {
    public:
        F64 mass;
        F64vec pos;
        F64mat quad;
        F64ort vertex_out_;
        F64ort vertex_in_;
    };
}
```

• Name of the class PS::MomentQuadrupoleScatter

• Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

quad: accumulated quadrupole.

vertex\_out\_: positions of vertices of the smallest rectangular parallelepiped including all grouped particles when each particle is regarded as a sphere whose radius is the returned value of EssentialParticleJ::getRSearch.

vertex\_in\_: positions of vertices of the smallest rectangular parallelepiped including all grouped particles.

• Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge, EssentialParticleJ::getPos and EssentialParticleJ::getRSearch and returns mass/electron charge, position and cutoff length. Users can use an arbitrary name instead of the name of member functions of EssentialParticleJ.

# 7.5.2.4 PS::SEARCH\_MODE\_LONG\_SYMMETRY

# 7.5.2.4.1 PS::MomentMonopoleSymmetry

This class encapsulates the monopole moment. The reference point for the calculation of monopole is set to center of mass or center of charge.

```
namespace ParticleSimulator {
   class MomentMonopoleSymmetry {
    public:
        F64 mass;
        F64vec pos;
        F64ort vertex_out_;
        F64ort vertex_in_;
    };
}
```

- Name of the class PS::MomentMonopoleSymmetry
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

vertex\_out\_: positions of vertices of the smallest rectangular parallelepiped including all grouped particles when each particle is regarded as a sphere whose radius is the returned value of EssentialParticleJ::getRSearch.

vertex\_in\_: positions of vertices of the smallest rectangular parallelepiped including all grouped particles.

#### • Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge, EssentialParticleJ::getPos and EssentialParticleJ::getRSearch and returns mass/electron charge, position and cutoff length. Users can use an arbitrary name instead of the name of member functions of EssentialParticleJ.

# 7.5.2.4.2 PS::MomentQuadrupoleSymmetry

This class encapsulates the monopole and quadrupole moments. The reference point for the calculation of monopole is set to center of mass or center of charge.

```
namespace ParticleSimulator {
   class MomentQuadrupoleSymmetry {
    public:
        F64       mass;
        F64vec pos;
        F64mat quad;
        F64ort vertex_out_;
        F64ort vertex_in_;
    };
}
```

- Name of the class PS::MomentQuadrupoleSymmetry
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

quad: accumulated quadrupole.

vertex\_out\_: positions of vertices of the smallest rectangular parallelepiped including all grouped particles when each particle is regarded as a sphere whose radius is the returned value of EssentialParticleJ::getRSearch.

vertex\_in\_: positions of vertices of the smallest rectangular parallelepiped including all grouped particles.

#### • Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge, EssentialParticleJ::getPos and EssentialParticleJ::getRSearch and returns mass/electron charge, position and cutoff length. Users can use an arbitrary name instead of the name of member functions of EssentialParticleJ.

# 7.5.2.5 PS::SEARCH\_MODE\_LONG\_CUTOFF

# 7.5.2.5.1 PS::MomentMonopoleCutoff

This class encapsulates the monopole moment. The reference point for the calculation of monopole is set to center of mass or center of charge.

```
namespace ParticleSimulator {
    class MomentMonopoleCutoff {
    public:
        F64 mass;
        F64vec pos;
    };
}
```

- Name of the class PS::MomentMonopoleCutoff
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

• Terms of use

The class EssentialParticleJ (see, Sec. 7.4) has EssentialParticleJ::getCharge, EssentialParticleJ::getPos and EssentialParticleJ::getRSearch and returns mass/electron charge, position and cutoff length. Users can use an arbitrary name instead of the name of member functions of EssentialParticleJ.

# 7.5.3 Required member functions

# 7.5.3.1 Summary

Below, the required member functions of Moment class are described. In this section we use the name Mom as a Moment class.

#### 7.5.3.2 Constructor

```
class Mom {
public:
    Mom ();
};
```

• Arguments

None.

• Returns

None.

# • Behaviour

Initialize the instance of Mom class.

# 7.5.3.3 Mom::init

```
class Mom {
public:
    void init();
};
```

# • Arguments

None.

# • Returns

None.

# • Behaviour

Initialize the instance of Mom class.

# 7.5.3.4 Mom::getPos

```
class Mom {
public:
    PS::F32vec getPos();
};
```

# • Arguments

None.

# • Returns

PS::F32vec or PS::F64vec. Returns the position of a variable of class.

# 7.5.3.5 Mom::getCharge

```
class Mom {
public:
    PS::F32 getCharge() const;
};
```

# • Arguments

None.

# • Returns

PS::F32 or PS::F64. Returns the mass or charge of a variable of class Mom.

# 7.5.3.6 Mom::accumulateAtLeaf

```
class Mom {
public:
    template <class Tepj>
    void accumulateAtLeaf(const Tepj & epj);
};
```

# • Arguments

```
epj: Input. const Tepj & type. The object of Tepj.
```

# • Returns

None.

# • Behaviour

Accumulate the multipole moment from EssentialParticleJ.

# 7.5.3.7 Mom::accumulate

```
class Mom {
  public:
     void accumulate(const Mom & mom);
};
```

# • Arguments

mom: Input. const Mom & type. The instance of Mom class.

#### • Returns

None.

# • Behaviour

Accumulate the multipole moments in Mom class from Mom class objects.

## 7.5.3.8 Mom::set

```
class Mom {
public:
    void set();
};
```

None.

#### • Returns

None.

#### • Behaviour

Normalize the multipole moments, since the member functions Mom::accumulateAtLeaf and Mom::accumulate do not normalize the multipole moment.

# 7.5.3.9 Mom::accumulateAtLeaf2

```
class Mom {
public:
    template <class Tepj>
    void accumulateAtLeaf2(const Tepj & epj);
};
```

# • Arguments

epj: Input. const Tepj & type. The instance of Teps.

### • Returns

None.

# • Behaviour

Accumulate the multipole moments in Mom class from EssentialParticleJ class objects.

#### 7.5.3.10 Mom::accumulate2

```
class Mom {
public:
    void accumulate(const Mom & mom);
};
```

# • Arguments

mom: Input. const Mom & type. The instance of Mom class.

# • Returns

None.

#### • Behaviour

Accumulate the multipole moments in Mom class from Mom class objects.

# 7.5.4 Required member functions for specific cases

# 7.5.4.1 Summary

Below, the required member functions of Moment class for specific cases are described. In this section we use the name Mom as a Moment class.

# 7.5.4.2 One of PS::SEARCH\_MODE\_LONG\_CUTOFF,

PS::SEARCH\_MODE\_LONG\_SCATTER,
PS::SEARCH\_MODE\_LONG\_SYMMETRY
is used as PS::SEARCH\_MODE

# 7.5.4.3 Mom::getVertexIn

```
class Mom {
public:
    F64ort getVertexIn();
};
```

### • Arguments

None.

#### • Returns

PS::F32ort or PS::F64ort types.

### • Behaviour

Return the positions of two vertices describing the smallest cuboid (rectangle if the space dimension is two) that contains all the particles corresponding to this Mom class.

#### • Notes

The positional information above must be calculated correctly in each tree cell. For this end, users must implement coordinate calculations into the member functions accumulateAtLeaf and accumulate, which are used, respectively, to calculate the moment information at the leaf cells (i.e. the tree cells at the deepest levels of a tree structure) from particles and to calculate the moment information of a parent tree cell from its child tree cells. Below, we show an example of the implementation of the member function getVertexIn.

```
class Mom {
public:
    // vertex_in_: Member variable storing the positional
                   information of a cuboid (or rectangle)
    F64ort vertex_in_;
    F64ort getVertexIn() const { return vertex_in_; }
    template < class Tep; >
    void accumulateAtLeaf(const Tepj & epj){
         // Other calculations
        (this->vertex_in_).merge(epj.getPos());
    }
    void accumulate(const Mom & mom){
        // Other calculations
        (this->vertex_in_).merge(mom.vertex_in_);
    }
};
```

where merge is one of the member functions of Orthotope type. Users can freely change the name of this member variable.

# 7.5.4.4 Mom::getVertexOut

```
class Mom {
public:
    F64ort getVertexOut();
};
```

# • Arguments

None.

#### • Returns

PS::F32ort or PS::F64ort type.

#### • Behaviour

Return the positions of two vertices describing the smallest cuboid (rectangle if the space dimension is two) that contains all of spheres (circles if the space dimension is two) whose centers and radii are, respectively, the positions and the returned values of getRSearch of particles corresponding to this Mom class.

# • Notes

As in the case of member function <code>getVertexIn</code>, the positional information above must be calculated correctly in each tree cell. For this end, users must implement coordinate calculations in the member functions <code>accumulateAtLeaf</code> and <code>accumulate.Below</code>, we show an example of the implementation of the member function <code>getVertexOut</code>.

```
class Mom {
public:
    // vertex_out_: Member variable storing the positional
                    information of a cuboid (or rectangle)
    F64ort vertex_out_;
    F64ort getVertexOut() const { return vertex_out_; }
    template < class Tep; >
    void accumulateAtLeaf(const Tepj & epj){
         // Other calculations
         (this->vertex_out_).merge(epj.getPos(), epj.getRSearch());
    }
    void accumulate(const Mom & mom){
        // Other calculations
        (this->vertex_out_).merge(mom.vertex_out_);
    }
};
```

where merge is a member function of Orthotope type. Users can freely change the name of the member variable.

# 7.6 SuperParticleJ

# 7.6.1 Summary

The class SuperParticleJ encapsulates the informations of a grouped particles (see, step 0 in Sec. 2.3). This class is required if one of the followings are employed as PS::SEARCH\_MODE.

- PS::SEARCH\_MODE\_LONG
- PS::SEARCH\_MODE\_LONG\_SCATTER
- PS::SEARCH\_MODE\_LONG\_SYMMETRY
- PS::SEARCH\_MODE\_LONG\_CUTOFF

This class has a member function which is used for data-transfer between FDPS. This class is related to Moment class. Thus, this class has member functions which cast the Moment class to this class (and vice versa).

Similar to Moment class, since this class has prescribed rules, FDPS has pre-defined classes. This section aims to describe the specific member functions both always and in specific case required for SuperParticleJ.

#### 7.6.2 Pre-defined class

FDPS has some pre-defined SuperParticleJ class. Below, the available classes for each PS::SEARCH\_MODE are described.

# 7.6.2.1 PS::SEARCH\_MODE\_LONG

# 7.6.2.1.1 *PS::SPJMonopole*

A SuperParticleJ class which is related to PS::MomentMonopole class. It contains up to monopole moment.

```
namespace ParticleSimulator {
   class SPJMonopole {
    public:
        F64 mass;
        F64vec pos;
    };
}
```

- Name of the class PS::SPJMonopole
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

• Terms of use

The same as PS::MomentMonopole class.

# 7.6.2.1.2 PS::SPJQuadrupole

A SuperParticleJ class which is related to PS::MomentQuadrupole class. It contains up to quadrupole moment.

```
namespace ParticleSimulator {
   class SPJQuadrupole {
    public:
        F64 mass;
        F64vec pos;
        F64mat quad;
    };
}
```

- Name of the class PS::SPJQuadrupole
- Members and their information

mass: accumulated mass.

pos: center of mass.

quad: accumulated quadrupole.

• Terms of use

The same as PS::MomentQuadrupole class.

# 7.6.2.1.3 PS::SPJMonopoleGeometricCenter

A SuperParticleJ class which is related to PS::MomentMonopoleGeometricCenter class. It contains up to quadrupole moment (the reference point is set to geometric center of grouped particles).

- Name of the class PS::SPJMonopoleGeometricCenter
- Members and their information

charge: accumulated mass/charge.

pos: geometric center.

• Terms of use

The same as PS::MomentMonopoleGeometricCenter class.

# 7.6.2.1.4 PS::SPJDipoleGeometricCenter

A SuperParticleJ class which is related to PS::MomentDipoleGeometricCenter class. It contains up to dipole moment (the reference point is set to geometric center of grouped particles).

- Name of the class PS::SPJDipoleGeometricCenter
- Members and their information

charge: accumulated mass/charge.

pos: geometric center.

dipole: dipole of the particle mass or electron charge.

• Terms of use

The same as PS::MomentDipoleGeometricCenter class.

# 7.6.2.1.5 PS::SPJQuadrupoleGeometricCenter

A SuperParticleJ class which is related to PS::MomentQuadrupoleGeometricCenter class. It contains up to quadrupole moment (the reference point is set to geometric center of grouped particles).

- Name of the class PS::SPJQuadrupoleGeometricCenter
- Members and their information

charge: accumulated mass/charge.

pos: geometric center.

dipole: dipole of the particle mass or electron charge.

quadrupole: quadrupole of the particle mass or electron charge.

• Terms of use

The same as PS::MomentQuadrupoleGeometricCenter class.

# 7.6.2.2 PS::SEARCH\_MODE\_LONG\_SCATTER

# 7.6.2.2.1 PS::SPJMonopoleScatter

A SuperParticleJ class which is related to PS::MomentMonopoleScatter class. It contains up to monopole moment.

```
namespace ParticleSimulator {
    class SPJMonopoleScatter {
    public:
        F64 mass;
        F64vec pos;
    };
}
```

- Name of the class PS::SPJMonopoleScatter
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

• Terms of use

The same as PS::MomentMonopoleScatter class.

# 7.6.2.2.2 PS::SPJQuadrupoleScatter

A SuperParticleJ class which is related to PS::MomentQuadrupoleScatter class. It contains up to quadrupole moment.

```
namespace ParticleSimulator {
   class SPJQuadrupoleScatter {
    public:
        F64 mass;
        F64vec pos;
        F64mat quad;
    };
}
```

- Name of the class PS::SPJQuadpoleScatter
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

quad: accumulated quadrupole.

• Terms of use

The same as PS::MomentQuadrupoleScatter class.

# 7.6.2.3 PS::SEARCH\_MODE\_LONG\_SYMMETRY

# 7.6.2.3.1 PS::SPJMonopoleSymmetry

A SuperParticleJ class which is related to PS::MomentMonopoleSymmetry class. It contains up to monopole moment.

```
namespace ParticleSimulator {
   class SPJMonopoleSymmetry {
    public:
        F64 mass;
        F64vec pos;
   };
}
```

- Name of the class PS::SPJMonopoleSymmetry
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

• Terms of use

The same as PS::MomentMonopoleSymmetry class.

# 7.6.2.3.2 PS::SPJQuadrupoleSymmetry

A SuperParticleJ class which is related to PS::MomentQuadrupoleSymmetry class. It contains up to quadrupole moment.

```
namespace ParticleSimulator {
   class SPJQuadrupoleSymmetry {
    public:
        F64 mass;
        F64vec pos;
        F64mat quad;
    };
}
```

- Name of the class PS::SPJQuadpoleSymmetry
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

quad: accumulated quadrupole.

Terms of use

The same as PS::MomentQuadrupoleSymmetry class.

# 7.6.2.4 PS::SEARCH\_MODE\_LONG\_CUTOFF

# 7.6.2.4.1 PS::SPJMonopoleCutoff

A SuperParticleJ class which is related to PS::MomentMonopoleCutoff class. It contains up to monopole moment.

```
namespace ParticleSimulator {
    class SPJMonopoleCutoff {
    public:
        F64 mass;
        F64vec pos;
    };
}
```

- Name of the class PS::SPJMonopoleCutoff
- Members and their information

mass: accumulated mass or electron charge.

pos: center of mass or center of charge.

• Terms of use

The same as PS::MomentMonopoleCutoff class.

# 7.6.3 Required member functions

# 7.6.3.1 Summary

Below, the required member functions of SuperParticleJ class are described. In this section we use the name SPJ as a SuperParticleJ class.

# 7.6.3.2 SPJ::getPos

```
class SPJ {
public:
    PS::F64vec getPos() const;
};
```

# • Arguments

None.

#### • Returns

PS::F32vec or PS::F64vec. Returns the position pf a super-particle of class SPJ.

# 7.6.3.3 SPJ::setPos

```
class SPJ {
  public:
    void setPos(const PS::F64vec pos_new);
};
```

# • Arguments

pos\_new: Input. const PS::F32vec or const PS::F64vec. Modified positions of particle by FDPS.

## • Returns

None.

# • Behaviour

Replaces the positions in SPJ class by those modified by FDPS.

# 7.6.3.4 SPJ::copyFromMoment

```
class Mom;
class SPJ {
  public:
    void copyFromMoment(const Mom & mom);
};
```

# • Arguments

 ${\tt mom:}$  Input. const  ${\tt Mom}$  & type.  ${\tt Mom}$  can be both user defined and pre-defined  ${\tt Moment}$  class.

#### • Returns

None.

#### • Behaviour

Copies the informations of Mom class to SPJ.

# 7.6.3.5 SPJ::convertToMoment

# • Arguments

None.

#### • Returns

Mom type. The constructor of Mom class.

### • Behaviour

Returns the constructor of Mom class.

# 7.6.3.6 SPJ::clear

```
class SPJ {
  public:
    void clear();
};
```

# • Arguments

None.

#### • Returns

None.

# • Behaviour

Clears the information of SPJ class.

# 7.7 Force

# **7.7.1** Summary

The Force class contains the results of the calculation of interactions (see, step 0 in Sec. 2.3). In this section we describe the member functions in any case required.

#### 7.7.2 Premise

Let us take Result class as an example of Force as below. Users can use an arbitrary name instead of Result.

# 7.7.3 Required member functions

The member function Result::clear is in any case needed. This function initialize the result on interaction calculations.

#### 7.7.3.1 Result::clear

```
class Result {
  public:
    void clear();
};
```

# • Arguments

None.

#### • Returns

None.

#### • Behaviour

Initializes the result of interaction calculations.

# 7.8 Header

# 7.8.1 Summary

Header class defines the format of header of input/output files. This class is required if users use API for file I/O provided by FDPS (ParticleSystem::readParticleAscii, ParticleSystem::readParticleBinary, and ParticleSystem and want to add header information to input/output file. Below, we describe the rules for defining the member functions required from provided input/output API. There is no functions which is always required.

### 7.8.2 Premise

Let us take Hdr class as an example of Header as below. Users can use an arbitrary name in place of Hdr.

# 7.8.3 Required member functions for specific case

# 7.8.3.1 Hdr::readAscii

```
class Hdr {
public:
    PS::S32 readAscii(FILE *fp);
};
```

# • Arguments

fp: Input. FILE \* type. The file pointer of the input file.

# • Returns

PS::S32 type. Returns the value of number of particles. Returns -1 if header does not contain the number of particle.

# • Behavior

Read the header data from the input file.

#### 7.8.3.2 Hdr::writeAscii

```
class Hdr {
public:
    void writeAscii(FILE *fp);
};
```

#### • Arguments

fp: Input. FILE \* type. The file pointer of the output file.

#### • Returns

None.

# • Behavior

Copies the header information to the output file.

# 7.8.3.3 Hdr::readBinary

```
class Hdr {
public:
    PS::S32 readBinary(FILE *fp);
};
```

fp: Input. FILE \* type. The file pointer of the input file.

#### • Returns

PS::S32 type. Returns the value of number of particles. Returns -1 if header does not contain the number of particle.

#### • Behavior

Read the header data from the input file.

# 7.8.3.4 Hdr::writeBinary

```
class Hdr {
public:
    void writeBinary(FILE *fp);
};
```

# • Arguments

fp: Input. FILE \* type. The file pointer of the output file.

#### • Returns

None.

#### • Behavior

Copies the header information to the output file.

# 7.9 Functor calcForceEpEp

# 7.9.1 Summary

Functor calcForceEpEp defines the interaction between two particles. This functor is required for the calculation of interactions (see, step 0 in Sec. 2.3).

### 7.9.2 Premise

Here one example of gravitational *N*-body problems are shown. The name of functor calcForceEpEp is gravityEpEp, which is an arbitrary. The class name of EssentialParitlceI, EssentialParitlceJ and Force are EPI, EPJ and Result.

# 7.9.3 gravityEpEp::operator ()

# Listing 19: calcForceEpEp

```
1 class Result;
```

```
class EPJ;
3
  struct gravityEpEp {
5
       static PS::F32 eps2;
6
       void operator () (const EPI *epi,
7
                          const PS::S32 ni,
8
                          const EPJ *epj,
9
                          const PS::S32 nj,
10
                          Result *result)
       }
11
12 };
13 PS::F32 gravityEpEp::eps2 = 9.765625e-4;
```

```
epi: Input. const EPI * type or EPI * type. Array of i particles.
ni: Input. const PS::S32 type or PS::S32. The number of i particles.
epj: Input. const EPJ * type or EPJ * type. Array of j particles.
nj: Input. const PS::S32 type or PS::S32 type. Number of j particles.
result: Output. Result * type. Array of the results of interaction.
```

#### • Returns

None.

### • Behaviour

Calculates the interaction to i- particle from j- particle.

# 7.10 Functor calcForceSpEp

#### **7.10.1** Summary

Functor calcForceSpEp defines the interaction to a particle from super particle. This functor is required for the calculation of interactions (see, step 0 in Sec. 2.3).

### 7.10.2 Premise

Here one example of gravitational *N*-body problems are shown. The superparticles are regarded to be constructed from up to monopole. The name of functor calcForceSpEp is gravitySpEp, which is an arbitrary. The class name of EssentialParitlceI, SuperParitlceJ and Force are EPI, SPJ and Result.

# 7.10.3 gravitySpEp::operator ()

```
Listing 20: calcForceSpEp
```

```
1 class Result;
2 class EPI;
```

```
3
  class SPJ;
   struct gravitySpEp {
5
       static PS::F32 eps2;
6
       void operator () (const EPI *epi,
7
                           const PS::S32 ni,
8
                           const SPJ *spj,
9
                           const PS::S32 nj,
10
                           Result *result);
11 };
```

```
epi: Input. const EPI * type or EPI * type. Array of i particles.
ni: Input. const PS::S32 type or PS::S32. The number of i particles.
spj: Input. const SPJ * type or SPJ * type. Array of super particle.
nj: Input. const PS::S32 type or PS::S32 type. Number of super particles.
result: Output. Result * type. Array of the results of interaction.
```

#### • Returns

None.

#### Behaviour

Calculate interactions from super particle to *i*-th particle.

# 7.11 Functor calcForceDispatch

#### **7.11.1** Summary

Functor calcForceDispatch, paired with functor calcForceRetrieve, defines the interaction between two particles. This functor can be used for the calculation of interactions (see, step 0 in Sec. 2.3), instead of calcForceSpEp and calcForceEpEp. The difference from calcForceSpEp and calcForceEpEp is that calcForceDispatch receives multiple interactions lists and i-particle lists. By doing so, it reduces the numbere of kernel calls to acclerators such as GPGPU, and improves the efficiency. The name of the functor calcForceDispatch is GravityDispatch. The name for classes EssentialParitlceI, EssentialParitlceJ and SuperParitlceJ are EPI, EPJ, and SPI, respectively. These names can be changed to any name legal.

# 7.11.2 The case of short-range interactions

Listing 21: calcForceDispatch

```
1 class EPI;
2 class EPJ;
3 PS::S32 HydroforceDispatch(const PS::S32 tag,
4 const PS::S32 nwalk,
```

tag: input. Type const PS::S3. The value should be non-negative and less than the value specified by the third argument of function

PS::TreeForForce::calcForceAllandWriteBackMultiWalk(). Corresponding call to CalcForceRetrieve() should use the same value for tag.

nwalk: input.Type const PS::S32. The number of interaction lists. The value should
not exceed the value of the sixth argument of

PS::TreeForForce::calcForceAllandWriteBackMultiWalk().

epi: input. Type const EPI\*\*. Array of the array of i-particles.

ni: input. Type const PS::S32\*. Array of the numbers of i-particles.

epj: input. Type const EPJ\*\*. Array of the array of j-particles.

nj: input. Type const PS::S32\*. Array of the number of j-paricles.

### • Return value

Type PS::S32. Should return 0 upon normal completion, and otherwize non-zero value.

# • Function

Send epi and epj to the accelerator and let the accelerator do the interaction calculation.

# 7.11.3 Case of long-range interaction

Listing 22: calcForceDispatch

```
1 class EPI;
2 class EPJ;
3 class SPJ;
  PS::S32 GravityDispatch(const PS::S32
                                                tag,
5
                              const PS::S32
                                                nwalk,
6
                              const EPI**
                                                epi,
7
                              const PS::S32*
                                                ni,
8
                              const EPJ**
                                                epj,
9
                              const PS::S32*
                                                nj_ep,
10
                              const SPJ**
                                                spj,
11
                              const PS::S32*
                                                nj_sp);
12 };
```

tag: input. Type const PS::S3. The value should be non-negative and less than the value specified by the third argument of function

PS::TreeForForce::calcForceAllandWriteBackMultiWalk(). Corresponding call to CalcForceRetrieve() should use the same value for tag.

nwalk: input.Type const PS::S32. The number of interaction lists. The value should
not exceed the value of the sixth argument of

PS::TreeForForce::calcForceAllandWriteBackMultiWalk().

epi: input. Type const EPI\*\*. Array of the array of i-particles.

ni: input. Type const PS::S32\*. Array of the numbers of i-particles.

epj: input. Type const EPJ\*\*. Array of the array of j-particles.

nj: input. Type const PS::S32\*. Array of the number of j-paricles.

spj: input. Type const SPJ\*\*. Array of the array of superparticles.

nj\_sp: input. Type const PS::S32\*. Array of the number of superparicles.

# • Return value

Type PS::S32. Should return 0 upon normal completion, and otherwize non-zero value.

#### • Function

Send epi and epj to the accelerator and let the accelerator do the interaction calculation.

# 7.12 Functor calcForceRetrieve

#### **7.12.1** Summary

Functor calcForceRetrieve retrieves the result of interaction calculation dispatched by functor calcForceDispatch.

The name of functor calcForceRetrieve is GravityRetrieve and the name of Force class is Result

### Listing 23: calcForceDispatch

```
1 class EPI;
2 class EPJ;
3 class Result;
4 PS::S32 GravityRetrieve(const PS::S32
                                             tag,
5
                            const PS::S32
                                             nwalk,
6
                            const PS::S32
                                                     [],
                                             ni
7
                                             result [][]);
                            Result
8 };
```

tag: input. Type const PS::S32. Should be the same as that for the corresponding call to CalcForceDispatch().

nwalk: input. Type const PS::S32. The number of interaction lists. Should be the same as that for the corresponding call to CalcForceDispatch().

ni: input. Type const PS::S32\* or PS::S32\*. Array of the numbers of i-particles.
result: output. Type Result\*\*.

• return value Returns 0 upon normal completion. Otherwize non-zero values are returned.

#### • Function

Store the results calculated by the call to CalcForceDispatch() with the same tag value to the array force.

# 8 Initialization and Finalization of FDPS

### 8.1 General

This section describes the APIs of initialization and finalization of FDPS.

# 8.2 API

### 8.2.1 PS::Initialize

This API must be called to initialize FDPS.

# • arguments

```
argo: input. Type PS::S32 &. The total number of arguments on the command line. argv: input. Type char ** &. The pointer to pointer to strings on the command line.
```

# • returned value

void.

# • function

Initialize FDPS. This API must be called before other APIs of FDPS are called. Note, since MPI::Init is called, that inside this API arguments "argc" and "argv" may change.

# 8.2.2 PS::Finalize

To finalize FDPS, this API must be called.

```
void PS::Finalize();
```

#### • arguments

void.

#### • return value

void.

# • function

Finalize FDPS.

# 8.2.3 PS::Abort

```
void PS::Abort(const PS::S32 err = -1);
```

# • arguments

err: Input. Type const PS::S32. The termination status of a program. Default value is -1.

# • returned value

void.

# • function

Terminate the user program abnormally. The argument is the termination status of the program. Under MPI environment, MPI::Abort() is called, otherwise exit() is called.

# 8.2.4 PS::DisplayInfo

```
void PS::DisplayInfo();
```

# • arguments

void.

# • returned value

void.

# • function

Show the license and other information of FDPS.

# 9 Classes defined in FDPS and their APIs

This section describes the classes defined in FDPS and their APIs. The following subsections describe the standard classes and the extention classes of FDPS.

# 9.1 Standard Classes

# 9.1.1 Summary

The following subsections describe the four standard classes of FDPS: DomainInfo class, ParticleSystem class, TreeForForce class and Comm class.

### 9.1.2 DomainInfo Class

This section describes DomainInfo class. This class handles the decomposition of computational domains and keeps their data.

# 9.1.2.1 Creation of Object

DomainInfo class is declared as below.

# Listing 24: DomainInfo0

```
1 namespace ParticleSimulator {
2    class DomainInfo;
3 }
```

Next example shows how to create an object of DomainInfo class. Here, the object is named "dinfo".

```
PS::DomainInfo dinfo;
```

### 9.1.2.2 API

DomainInfo class has the APIs for initialization and decomposition of domain. The following subsections describe them.

#### 9.1.2.2.1 Initial Setup

The APIs for the initial setup of DomainInfo class are declared below.

### Listing 25: DomainInfo1

```
1 namespace ParticleSimulator {
2    class DomainInfo{
3     public:
4         DomainInfo();
5         void initialize(const F32 coef_ema=1.0);
6         void setNumberOfDomainMultiDimension(const S32 nx,
```

```
const S32 ny,
const S32 nz=1);

void setBoundaryCondition(enum BOUNDARY_CONDITION bc);

S32 getBoundaryCondition();

void setPosRootDomain(const F32vec & low,
const F32vec & high);

const F32vec & high);

};
```

### 9.1.2.2.1.1 Constructor

Constructor

```
void PS::DomainInfo::DomainInfo();
```

# • arguments

void.

#### • returned value

void.

#### • function

Create an object of DomainInfo class.

# 9.1.2.2.1.2 PS::DomainInfo::initialize

PS::DomainInfo::initialize

```
void PS::DomainInfo::initialize(const PS::F32 coef_ema=1.0);
```

#### • arguments

 $\mathtt{coef\_ema}$ : input. Type const PS::F32. The smoothing factor of an exponential moving average.

# • returned value

void.

#### function

Initialize an object of the domain information class. The argument coef\_ema is the smoothing factor of exponential moving average and is a constant real value between 0 and 1. If other values are chosen, FDPS sends an error message and terminates the user program. A larger coef\_ema weighs newer values rather than older values. In the case of unity, the domains are determined by using the newest values only and in the case of zero, they are determined by using the initial values only. Users call this

API only once. The details of this function are described in the paper by Ishiyama, Fukushige & Makino (2009, Publications of the Astronomical Society of Japan, 61, 1319)

# 9.1.2.2.1.3 PS::DomainInfo::setNumberOfDomainMultiDimension

PS::DomainInfo::setNumberOfDomainMultiDimension

# • arguments

nx: Input. Type const PS::S32. The number of subdomains along x direction.

ny: Input. Type const PS::S32. The number of subdomains along y direction.

nz: Input. Type const PS::S32. The number of subdomains along z direction. The default value is 1.

#### • returned value

void.

#### function

Set the numbers of subdomains. If the API is not called: nx, ny and nz are determined automatically. If the product of nx, ny and nz is not equal to the total number of MPI processes, FDPS sends an error message and terminates the user program.

# 9.1.2.2.1.4 PS::DomainInfo::setBoundaryCondition

PS::DomainInfo::setBoundaryCondition

#### • arguments

bc: input. Type enum PS::BOUNDARY\_CONDITION. Boundary conditions.

#### • returned value

void.

#### • function

Set the boundary condition. FDPS allows boundary conditions defined in Section 6.8.2 (BOUNDARY\_CONDITION\_SHEARING\_BOX and BOUNDARY\_CONDITION\_USER\_DEFINED have not been implemented yet). If the API is not called, the open boundary is used.

# 9.1.2.2.1.5 PS::DomainInfo::getBoundaryCondition

PS::DomainInfo::getBoundaryCondition

```
S32 PS::DomainInfo::getBoundaryCondition();
```

# • arguments

void.

# • returned value

Type PS::S32.

### • function

Return an integer value corresponding to the current boundary condition. The possible values are integers corresponding to the enumerators of PS::BOUNDARY\_CONDITION.

# 9.1.2.2.1.6 PS::DomainInfo::setPosRootDomain

PS::DomainInfo::setPosRootDomain

# • arguments

```
low: input. Type const PS::F32vec. Top vertex of the boundary (inclusive). high: input. Type const PS::F32vec. Bottom vertex of the boundary (exclusive).
```

### • returned value

void.

### function

Set positions of vertexes of top and bottom of root domain. The API do not need to be called under open boundary condition. Every coordinate of high must be greater than the corresponding coordinate of low. Otherwise, FDPS sends a error message and terminates the user program.

# 9.1.2.2.2 Decomposition of Domain

The APIs of decomposition of domain are declared below:

# Listing 26: DomainInfo2

```
1 namespace ParticleSimulator {
2     class DomainInfo{
3     public:
```

```
4
            template < class Tpsys >
5
            void collectSampleParticle(Tpsys & psys,
6
                                          const bool clear,
7
                                          const F32 weight);
8
            template < class Tpsys >
9
            void collectSampleParticle(Tpsys & psys,
10
                                          const bool clear);
            template < class Tpsys >
11
12
            void collectSampleParticle(Tpsys & psys);
13
14
            void decomposeDomain();
15
16
            template < class Tpsys >
17
            void decomposeDomainAll(Tpsys & psys,
18
                                      const F32 weight);
19
            template < class Tpsys >
20
            void decomposeDomainAll(Tpsys & psys);
21
       };
22 }
```

# $9.1.2.2.2.1 \quad PS::DomainInfo::collectSampleParticle$

PS::DomainInfo::collectSampleParticle

# • arguments

psys: input. Type ParticleSystem &. ParticleSystem class for giving sample particles to decompose domain.

clear: input. Type const bool. A flag to clear the data of sample particles, if set to

weight: input. Type const PS::F32. A weight to determine the number of sample particles.

## • returned value

void.

# • function

Sample particles from an object of ParticleSystem class. If clear is true, the data of the samples collected before is cleared. Larger weight leads to give more sample

particles. More specifically,  $n_{\rm smpl}n_{\rm proc}(w_i/\sum_k w_k)$  particles are sampled from process i, where  $n_{\rm smpl}$  is the number of sampled particles per process, which is set by API setAverateTargetNumberOfSampleParticlePerProcess,

 $n_{\text{proc}}$  is the number of MPI processes, and  $w_i$  is weight of process i. For better load balance, weight should be a quantity that reflects the calculation time of each process.

### • arguments

psys: input. Type ParticleSystem &. An object of ParticleSystem class for giving sample particles to decompose domain.

clear: input. Type const bool. A flag to clear the data of particles, if set to 'true'.

#### returned value

void.

### • function

Sample particles from psys. If clear is true, the data are cleared.

### • arguments

psys: input. Type ParticleSystem &. An object of ParticleSystem class for giving sample particles to decompose domain.

### • returned value

void.

# • functions

Sample particles from psys.

# 9.1.2.2.2.2 PS::DomainInfo::decomposeDomain

PS::DomainInfo::decomposeDomain

```
void PS::DomainInfo::decomposeDomain();
```

# • arguments

void.

### • returned value

void.

### • function

Decompose calculation domains.

# 9.1.2.2.2.3 PS::DomainInfo::decomposeDomainAll

PS::DomainInfo::decomposeDomainAll

## • arguments

psys: input. Type ParticleSystem &. The particle system class giving sample particles for the domain decomposition.

weight: input. Type const PS::F32. The sampling weight.

### • returned value

void.

## • function

Sample particles from psys and decompose domains. This API is the combination of PS::DomainInfo::collectSampleParticle and PS::DomainInfo::decomposeDomain.

# • arguments

psys: input. Type ParticleSystem &. The particle system class for giving samples.

# • returned value

void.

# • function

Sample particles from "psys" and decompose domains. This API is the combination of PS::DomainInfo::collectSampleParticle and PS::DomainInfo::decomposeDomain.

### 9.1.2.2.3 Time Measurment

The member functions of collecting information of an object are declared below. When a member function is called, its execution time is set to the private member of TimeProfile class time\_profile. Until the method clearTimeProfile() is called, execution times are accumulated.

Listing 27: DomainInfo3

```
1 namespace ParticleSimulator {
2    class DomainInfo{
3    public:
4         TimeProfile getTimeProfile();
5         void clearTimeProfile();
6    };
7 }
```

# 9.1.2.2.3.1 PS::DomainInfo::getTimeProfile

PS::DomainInfo::getTimeProfile

```
PS::TimeProfile PS::DomainInfo::getTimeProfile();
```

• arguments

void.

• returned value

Type PS::TimeProfile.

• function

Return the copy of the member variable time\_profile\_.

# 9.1.2.2.3.2 PS::DomainInfo::clearTimeProfile

PS::DomainInfo::clearTimeProfile

```
void PS::DomainInfo::clearTimeProfile();
```

• arguments

void.

• returned value

void.

• function

Set all member variables of time\_profile\_ to 0.

# 9.1.2.2.4 Obtain Information

The APIs of obtaining information is deleared bellow.

# Listing 28: DomainInfo3

# 9.1.2.2.4.1 PS::DomainInfo::getMemSizeUsed

PS:: Domain Info:: get Mem Size Used

```
PS::S64 PS::DomainInfo::getMemSizeUsed();
```

 $\bullet$  arguments

void.

• returned value

Type PS::S64. Size of used memory of the calling object in the unit of byte.

• function

Return the size of used memory of the calling object in the unit of byte.

# 9.1.3 ParticleSystem Class

This section describes ParticleSystem class. This class is used to keep the information of particles for the exchange of particles.

# 9.1.3.1 Creation of Object

The particle system class is declared below.

# Listing 29: ParticleSystem0

```
1 namespace ParticleSimulator {
2    template < class Tptcl >
3    class ParticleSystem;
4 }
```

The template argument is user-defined FullParticle class

Next example shows how to create an object of ParticleSystem class. Here, the object is named system.

```
PS::ParticleSystem<FP> system;
```

### 9.1.3.2 API

DomainInfo class has APIs for initialization, obtaining information, file I/O and exchanging particles. The following subsections describe them.

# 9.1.3.2.1 Initial Setup

The APIs of initialization are declared below.

Listing 30: ParticleSystem1

```
namespace ParticleSimulator {
1
2
       template < class Tptcl>
3
       class ParticleSystem{
4
       public:
5
           ParticleSystem();
6
           void initialize();
7
           void setAverageTargetNumberOfSampleParticlePerProcess
8
                             (const S32 & nsampleperprocess);
9
       };
10 }
```

# 9.1.3.2.1.1 Constructor

Constructor

```
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::ParticleSystem();
```

• arguments

void.

• returned value

void.

• function

Create an object of ParticleSystem class.

# 9.1.3.2.1.2 PS::ParticleSystem::initialize

PS::ParticleSystem::initialize

```
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::initialize();
```

# • arguments

void.

### • returned value

void.

### • function

Initialize an object of ParticleSystem class. Users must call this API once.

# 9.1.3.2.1.3 PS::ParticleSystem::

setAverageTargetNumberOfSampleParticlePerProcess

PS::ParticleSystem::

set Average Target Number Of Sample Particle Per Process

# • arguments

nsampleperprocess: input. Type const PS::S32 &. The average number of sample particles per MPI process.

### • returned value

void.

### function

Set the average number of sample particles per MPI process. If this function is not called, the average number is 30.

# 9.1.3.2.2 Obtain Information

The APIs of obtaining information are declared below.

# Listing 31: ParticleSystem2

```
1
  namespace ParticleSimulator {
2
       template < class Tptcl >
3
       class ParticleSystem{
4
       public:
5
           Tptcl & operator [] (const S32 id);
6
           S32 getNumberOfParticleLocal() const;
7
           S32 getNumberOfParticleGlobal() const;
8
           S64 getMemSizeUsed() const;
9
       };
10 }
```

# 9.1.3.2.2.1 PS::ParticleSystem::operator []

PS::ParticleSystem::operator []

## • arguments

id: input. Type const PS::S32. Index of the particle array in the calling object.

### • returned value

Type FullParticle &.

# • function

Return the reference of the id-th object of FullParticle class.

# 9.1.3.2.2.2 PS::ParticleSystem::getNumberOfParticleLocal

PS::ParticleSystem::getNumberOfParticleLocal

```
template <class Tptcl>
PS::S32 PS::ParticleSystem<Tptcl>::getNumberOfParticleLocal();
```

# • arguments

void.

### • returned value

Type PS::S32. The number of particles of the calling process.

# • function

Return the number of particles of the calling process.

# 9.1.3.2.2.3 PS::ParticleSystem::qetNumberOfParticleGlobal

PS:: Particle System:: get Number Of Particle Global

```
template <class Tptcl>
PS::S32 PS::ParticleSystem<Tptcl>::getNumberOfParticleGlobal();
```

# • arguments

void.

### • returned value

Type PS::S32. The total number of particles of all processes.

### • function

Return the total number of particles of all processes.

# 9.1.3.2.2.4 PS::DomainInfo::getUsedMemorySize

PS::DomainInfo::getUsedMemorySize

```
PS::S64 PS::DomainInfo::getUsedMemorySize();
```

### • arguments

void.

# • returned value

PS::S64. The size of used memory of the calling object in the unit of byte.

# • function

Return the size of used memory of the calling object in the unit of byte.

# 9.1.3.2.3 File I/O

The APIs of file I/O are declared below.

Listing 32: ParticleSystem3

```
namespace ParticleSimulator {
1
2
       template < class Tptcl >
3
       class ParticleSystem{
4
       public:
5
           template <class Theader>
6
           void readParticleAscii(const char * const filename,
7
                                    const char * const format,
8
                                    Theader & header);
9
           void readParticleAscii(const char * const filename,
10
                                    const char * const format);
11
           template <class Theader>
12
           void readParticleAscii(const char * const filename,
13
                                    Theader & header);
14
           void readParticleAscii(const char * const filename);
15
           template <class Theader>
           void readParticleAscii(const char * const filename,
16
17
                                    const char * const format,
18
                                    Theader & header,
                                    void (Tptcl::*pFunc)(FILE*));
19
20
           void readParticleAscii(const char * const filename,
21
                                    const char * const format,
22
                                   void (Tptcl::*pFunc)(FILE*));
```

```
23
           template <class Theader>
24
           void readParticleAscii(const char * const filename,
25
                                   Theader & header,
26
                                   void (Tptcl::*pFunc)(FILE*));
27
           void readParticleAscii(const char * const filename,
28
                                   void (Tptcl::*pFunc)(FILE*) );
29
30
           template <class Theader>
           void readParticleBinary(const char * const filename,
31
32
                                   const char * const format,
33
                                   Theader & header);
34
           void readParticleBinary(const char * const filename,
                                   const char * const format);
35
36
           template <class Theader>
37
           void readParticleBinary(const char * const filename,
38
                                   Theader & header);
39
           void readParticleBinary(const char * const filename);
40
           template <class Theader>
41
           void readParticleBinary(const char * const filename,
42
                                   const char * const format,
43
                                   Theader & header,
44
                                   void (Tptcl::*pFunc)(FILE*));
45
           void readParticleBinary(const char * const filename,
46
                                   const char * const format,
47
                                   void (Tptcl::*pFunc)(FILE*));
48
           template <class Theader>
49
           void readParticleBinary(const char * const filename,
50
                                   Theader & header,
51
                                   void (Tptcl::*pFunc)(FILE*));
52
           void readParticleBinary(const char * const filename,
53
                                   void (Tptcl::*pFunc)(FILE*) );
54
55
           template <class Theader>
           void writeParticleAscii(const char * const filename,
56
57
                                     const char * const format,
58
                                     const Theader & header);
59
           void writeParticleAscii(const char * const filename,
60
                                     const char * format);
61
           template <class Theader>
62
           void writeParticleAscii(const char * const filename,
                                    const Theader & header);
63
64
           void writeParticleAscii(const char * const filename);
65
           template <class Theader>
66
           void writeParticleAscii(const char * const filename,
67
                                    const char * const format,
68
                                    const Theader & header,
```

```
69
                                     void (Tptcl::*pFunc)(FILE*)
                                           const);
70
            void writeParticleAscii(const char * const filename,
71
                                      const char * format,
72.
                                     void (Tptcl::*pFunc)(FILE*)
                                           const);
73
            template <class Theader>
74
            void writeParticleAscii(const char * const filename,
75
                                      const Theader & header,
                                     void (Tptcl::*pFunc)(FILE*)
76
                                           const):
77
            void writeParticleAscii(const char * const filename,
                                     void (Tptcl::*pFunc)(FILE*)
78
                                           const);
79
80
            template <class Theader>
81
            void writeParticleBinary(const char * const filename,
82
                                     const char * const format,
83
                                     const Theader & header);
84
            void writeParticleBinary(const char * const filename,
85
                                      const char * format);
86
            template <class Theader>
87
            void writeParticleBinary(const char * const filename,
88
                                     const Theader & header);
89
            void writeParticleBinary(const char * const filename);
90
            template <class Theader>
            void writeParticleBinary(const char * const filename,
91
92
                                      const char * const format,
93
                                      const Theader & header,
94
                                     void (Tptcl::*pFunc)(FILE*)
                                           const);
95
            void writeParticleBinary(const char * const filename,
                                      const char * format,
96
97
                                     void (Tptcl::*pFunc)(FILE*)
                                           const);
98
            template <class Theader>
99
            void writeParticleBinary(const char * const filename,
100
                                     const Theader & header,
                                     void (Tptcl::*pFunc)(FILE*)
101
                                           const);
            void writeParticleBinary(const char * const filename,
102
                                      void (Tptcl::*pFunc)(FILE*)
103
                                           const);
104
       };
105 }
```

As describe above, there are four APIs for file I/O: readParticleAscii, readParticleBinary, writeParticleAscii, and writeParticleBinary. They are overloaded functions and each has 8 different interfaces, which results from existence or non-existence of the following arguments (the definitions of them will be described later).

- 1) template <class Theader> Theader & header
- 2) const char \* const format
- 3) void (Tptcl::\*pFunc)(FILE \*)

In the following, we describe each API. For brevity, we describe functions overloaded in a single section.

# 9.1.3.2.3.1 PS::ParticleSystem::readParticleAscii

PS::ParticleSystem::readParticleAscii

readParticleAscii has the following eight interfaces.

```
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptlc>::readParticleAscii
                 (const char * const filename,
                  const char * const format,
                  Theader & header);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::readParticleAscii
            (const char * const filename,
             const char * const format);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::readParticleAscii
            (const char * const filename,
             Theader & header):
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::readParticleAscii
            (const char * const filename);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptlc>::readParticleAscii
                 (const char * const filename,
                  const char * const format,
                  Theader & header,
                  void (Tptcl::*pFunc)(FILE*));
```

## • Arguments

filename: Input. Type const char \*. Prefix of name of an input file in the parallel read mode. Name of an input file in the single-process read mode.

format: Input. Type const char \*. Format of file name for parallel read mode.

header: Input. Type Theader &. Header information of a file.

pFunc: Input. Type void (Tptcl::\*)(FILE\*). Pointer to a member function of class Tptcl, which receives a FILE pointer and returns void.

# • Returned value

None.

# • Function

If argument format exists, each process reads data of particles from the input file specified by both filename and format and stores the data into FullParticle-type objects (parallel read mode). If argument format is not present, the root process reads data of particles from the input file named filename and distributes the data to other processes (single-process read mode). In both read modes, if argument header is not present, this API reads the file twice. The first one is to get the number of particles stored in the file. The second is to read particle data. Thus, this API estimates the number of particles assuming that data of one particle is stored in a single line.

In the parallel read mode, filename and format represent the prefix of name of an input file and the format of file name, respectively. Format specifier is the same as that of the standard C library. There must be three format specifiers. The first is string type and both the second and third are integer types. The first string is set to be filename. The second and third integers are the number of the total processes and

the rank id of the calling process, respectively. For instance, if filename is nbody, format is %s\_%03d\_%03d.init and the number of total process is 64, the process with rank id of 12 reads the file named nbody\_064\_012.init.

The API uses function pFunc to read data of one particle if argument pFunc exists. Otherwise, it uses member function readAscii of FullParticle class. Thus, users must define FullParticle::readAscii if required. The requirement specification and an example of implementation of readAscii are described in Sections 7.2.4.3.1 and A.1.4.3, respectively. Function pFunc must fulfill the same requirement specification as that of readAscii.

Member function readAscii of Theader class is used to read header information of the input file. Thus, uses must define Theader::readAscii. Its requirement specification and an example of implementation are described in Section 7.8.3.1 and A.7, respectively.

Files are opened in ascii mode.

# $9.1.3.2.3.2 \quad PS::Particle System:: read Particle Binary$

PS::ParticleSystem::readParticleBinary

readParticleBinary has the following eight interfaces.

```
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptlc>::readParticleBinary
                 (const char * const filename,
                  const char * const format,
                  Theader & header);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::readParticleBinary
            (const char * const filename,
             const char * const format);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::readParticleBinary
            (const char * const filename,
             Theader & header);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::readParticleBinary
            (const char * const filename);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptlc>::readParticleBinary
```

```
(const char * const filename,
                  const char * const format,
                  Theader & header,
                  void (Tptcl::*pFunc)(FILE*));
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::readParticleBinary
            (const char * const filename,
             const char * const format,
             void (Tptcl::*pFunc)(FILE*));
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::readParticleBinary
            (const char * const filename,
             Theader & header,
             void (Tptcl::*pFunc)(FILE*));
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::readParticleBinary
            (const char * const filename,
             void (Tptcl::*pFunc)(FILE*));
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::writeParticleAscii
            (const char * const filename,
             const Theader & header,
             void (Tptcl::*pFunc)(FILE*)const);
```

# • Arguments

filename: Input. Type const char \*. Prefix of the name of an input file in the parallel read mode. Name of an input file in the single-process read mode.

format: Input. Type const char \*. Format of file name in the parallel read mode.

header: Input. Type Theader &. Header information of a file.

pFunc: Input. Type void (Tptcl::\*)(FILE\*). Pointer to a member function of class Tptcl, which receives a FILE pointer and returns void.

### • Returned value

None.

### Function

It provides almost the same function as readParticleAscii, but there are the following differences.

- Files are opened in binary mode.
- The API estimates the number of particles stored in a file by examining the amount of change of the file position indicator before and after reading one particle actually. Hence, data of one particle does not need to end with a newline character (\n).

# $9.1.3.2.3.3 \quad PS::Particle System::write Particle Ascii$

PS::ParticleSystem::writeParticleAscii

writeParticleAscii has the following eight interfaces.

```
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::writeParticleAscii
            (const char * const filename,
             const char * const format,
             const Theader & header);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::writeParticleAscii
            (const char * const filename,
             const char * const format);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::writeParticleAscii
            (const char * const filename,
             const Theader & header);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::writeParticleAscii
            (const char * const filename);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::writeParticleAscii
            (const char * const filename,
             const char * const format,
             const Theader & header,
             void (Tptcl::*pFunc)(FILE*)const);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::writeParticleAscii
            (const char * const filename,
             const char * const format,
             void (Tptcl::*pFunc)(FILE*)const);
```

## • Arguments

filename: Input. Type const char \* const. Prefix of the name of an output file in the parallel write mode. Name of an output file in the single-process write mode.

format: Input. Type const char \* const. Format of file name in the parallel write mode.

header: Input. Type const Theader &. Header information of a file.

pFunc: Input. Type void (Tptcl::\*)(FILE\*)const. Pointer to a member function of class Tptcl, which receives a FILE pointer and returns void.

### • Returned value

None.

### • Function

If argument format exists, each process writes data of FullParticle-type objects to the output file specified by both filename and format (parallel write mode). If argument format is not present, the root process gathers data of FullParticle-type objects owned by other processes and then writes the collected data to the output file named filename (single-process write mode). In both modes, if argument header exists, header information is also output.

How to describe format is the same as the case of

PS::ParticleSystem::readParticleAscii.

The API uses function pFunc to write data of one particle if argument pFunc exists. Otherwise, it uses member function writeAscii of FullParticle class. Thus, users must define FullParticle::writeAscii if required. The requirement specification and an example of implementation of writeAscii are described in Sections 7.2.4.3.2 and A.1.4.3, respectively. Function pFunc must fulfill the same requirement specification as that of writeAscii.

Member function writeAscii of class Theader is used to write header information of the output file. Thus, users must define Theader::writeAscii. Its requirement

specification and an example of implementation are described in Sections 7.8.3.2 and A.7, respectively.

Files are opened in ascii mode.

# 9.1.3.2.3.4 PS::ParticleSystem::writeParticleBinary

PS::ParticleSystem::writeParticleBinary

writeParticleBinary has the following eight interfaces.

```
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::writeParticleBinary
            (const char * const filename,
             const char * const format,
             const Theader & header);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::writeParticleBinary
            (const char * const filename,
             const char * const format);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::writeParticleBinary
            (const char * const filename,
             const Theader & header);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::writeParticleBinary
            (const char * const filename);
template <class Tptcl>
template <class Theader>
void PS::ParticleSystem<Tptcl>::writeParticleBinary
            (const char * const filename,
             const char * const format,
             const Theader & header,
             void (Tptcl::*pFunc)(FILE*)const);
template <class Tptcl>
void PS::ParticleSystem<Tptcl>::writeParticleBinary
            (const char * const filename,
             const char * const format,
             void (Tptcl::*pFunc)(FILE*)const);
template <class Tptcl>
```

# • Arguments

filename: Input. Type const char \* const. Prefix of name of an input file in the parallel write mode. Name of an input file in the single-process write mode.

format: Input. Type const char \* const. Format of a file name in the parallel write mode.

header: Input. Type const Theader &. Header information of a file.

pFunc: Input. Type void (Tptcl::\*)(FILE\*)const. Pointer to a member function of class Tptcl, which receives a FILE pointer and returns void.

### • Returned value

None.

### • Function

It provide almost the same function as writeParticleAscii, but file are opened in binary mode in this API.

# 9.1.3.2.4 Exchange Particles

The APIs of exchange particle are declared below.

## Listing 33: ParticleSystem4

```
namespace ParticleSimulator {
   template < class Tptcl >
   class ParticleSystem {
    public:
        template < class Tdinfo >
        void exchangeParticle(Tdinfo & dinfo);
};
};
```

# 9.1.3.2.4.1 PS::ParticleSystem::exchangeParticle

PS::ParticleSystem::exchangeParticle

# • arguments

dinfo: input. Type DomainInfo &.

# • returned value

void.

### • function

Redistribute particles among MPI processes so that the particles are in appropriate domains.

# 9.1.3.2.5 Adding and removing particles

The APIs of adding and removing particles are declared as follows.

# 9.1.3.2.5.1 PS::ParticleSystem::addOneParticle()

PS::ParticleSystem::addOneParticle()

```
void PS::ParticleSystem::addOneParticle(const FullPartilce & fp);
```

### • arguments

fp: input. Type const FullParticle &. A particle to be added.

### • returned value

void.

# • function

Add a new particle fp at the end of the array of the particles.

# 9.1.3.2.5.2 PS::ParticleSystem::removeParticle()

PS::ParticleSystem::removeParticle()

# • arguments

idx: input. Type const PS::S32  $^{*}$  . An array of the index of the particles to be removed.

n: input. Type const PS::S32. The number of particles to be removed.

### • returned value

void.

### • function

Remove the particles with the index in the array idx. After calling this API, the order of the array of the particles would be changed.

### 9.1.3.2.6 Time Measurement

The APIs of Time Measurement are given below. When a member function is called, its execution time is set to the private member time\_profile. Until the method, clearTimeProfile(), execution times are accumulated.

Listing 34: ParticleSystem3

```
namespace ParticleSimulator {
1
2
       template < class Tptcl >
3
       class ParticleSystem{
4
       private:
5
            TimeProfile time_profile_;
6
       public:
7
            TimeProfile getTimeProfile();
8
            void clearTimeProfile();
9
10
       };
11 }
```

# 9.1.3.2.6.1 PS::ParticleSystem::qetTimeProfile

PS::ParticleSystem::getTimeProfile

```
PS::TimeProfile PS::ParticleSystem::getTimeProfile();
```

### • arguments

void.

### • returned value

Type PS::TimeProfile.

### • function

Return the copy of the member variable time\_profile\_.

# 9.1.3.2.6.2 PS::ParticleSystem::clearTimeProfile

PS::ParticleSystem::clearTimeProfile

```
void PS::ParticleSystem::clearTimeProfile();
```

• arguments

void.

• returned value

void.

• function

Set all member variables of time\_profile\_ to 0.

9.1.3.2.7 Others

Other APIs are declared below.

# Listing 35: ParticleSystem4

```
1 namespace ParticleSimulator {
2
       template < class Tptcl >
3
       class ParticleSystem{
4
       public:
5
           template < class Tdinfo >
6
           void adjustPositionIntoRootDomain
7
                         (const Tdinfo & dinfo);
8
           void setNumberOfParticleLocal(const S32 n);
9
           tamplate < class Tcomp >
           void sortParticle(Tcomp comp);
10
11
       };
12 }
```

# $9.1.3.2.7.1 \quad PS::Particle System:: adjust Position Into Root Domain$

PS:: Particle System:: adjust Position Into Root Domain

### • arguments

dinfo: input. Type const Tdinfo &. Object of the domain information class.

### • returned value

void.

### function

Under the periodic boundary condition, the particles outside the calculation domain move to appropriate positions.

# 9.1.3.2.7.2 PS::ParticleSystem::setNumberOfParticleLocal

PS::ParticleSystem::setNumberOfParticleLocal

### • arguments

n: input. Type const PS::S32. The number of particles of the calling process.

# • returned value

void.

### • function

Set the number of particles of the calling process.

# 9.1.3.2.7.3 PS::ParticleSystem::sortParticle

PS::ParticleSystem::sortParticle

```
tamplate<class Tcomp>
void sortParticle(Tcomp comp);
```

### • arguments

comp: input. Type Tcomp. Compare function which returns bool type value. This function needs two arguments.

The following is an example of comp to sort FullParticles in ascending order of particle ID.

```
bool comp(const FP & left, const FP & right){
    return left.id < right.id;
}</pre>
```

### • returned value

void.

### • function

This API sorts an array of FullParticles in the order determined by a comparison function comp. The returned value of comp must be bool type and it must take two arguments of FullParticles.

### 9.1.4 TreeForForce Class

This section describes TreeForForce class. This class is a module to calculate interactions between particles.

# 9.1.4.1 Creation of Object

This class is declared below.

Listing 36: TreeForForce0

```
namespace ParticleSimulator {
1
2
       template < class TSearchMode,
3
                 class TResult,
4
                 class TEpi,
5
                 class TEpj,
6
                 class TMomLocal,
7
                 class TMomGlobal,
8
                 class TSpj>
9
       class TreeForForce;
10 }
```

The types of template arguments are PS::SEARCH\_MODE class (user selected), Force class (user defined), EssentialParticleI class (user defined), EssentialParticleJ class (user defined), Moment class for local tree (user defined), Moment class for global tree (user defined) and SuperParticleJ class (user defined).

Depending on PS::SEARCH\_MODE, the wrapper classes to reduce the template arguments are available. Using these wrapper classes is highly recommended.

The following sections describe how to create objects by using the wrappers.

# 9.1.4.1.1 PS::SEARCH\_MODE\_LONG

This section describes the wrapper class for PS::SEARCH\_MODE\_LONG.

Next example shows how to create an object of PS::SEARCH\_MODE\_LONG class. Here, the object is named system.

```
PS::TreeForForceLong <TResult, TEpi, TEpj, TMom, TSpj>::Normal system;
```

The template arguments are Force class, EssentailParticleI class, EssentailParticleJ class, Moment class for both local and global trees and SuperParticleJ class. FDPS prepare some useful classes of the Moment and the SPJ (see section 7.5, 7.6). Users can use them.

More specialized wrappers are also prepared for some cases.

The next wrapper is for the case that the multipole expansion is up to the monopole and the expansion center is the barycenter of particles.

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::Monopole system;
```

The next wrapper is for the case that the multipole expansion is up to the quadrupole and the expansion center is the barycenter of particles.

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::Quadrupole system;
```

The next wrapper is for the case that the multipole expansion is up to the monopole and the expansion center is the geometric center of particles (not charge-weighted).

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::MonopoleGeometricCenter system;
```

The next wrapper is for the case that the multipole expansion is up to the dipole and the expansion center is the geometric center of particles (not charge-weighted).

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::DipoleGeometricCenter system;
```

The next wrapper is for the case that the multipole expansion is up to the quadrupole and the expansion center is the geometric center of particles (not charge-weighted).

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::QuadrupoleGeometricCenter system;
```

### 9.1.4.1.2 PS::SEARCH\_MODE\_LONG\_SCATTER

This section describes the wrapper class for PS::SEARCH\_MODE\_LONG\_SCATTER. Next example shows how to create an object of PS::SEARCH\_MODE\_LONG\_SCATTER class. Here, the object is named system.

```
PS::TreeForForceLong <TResult, TEpi, TEpj, TMom, TSpj>::WithScatterSearch system;
```

The template arguments are Force class, EssentialParticleI class, EssentialParticleJ class, Moment class for both local and global trees and SuperParticleJ class. FDPS prepare some useful classes of Moment and SuperParticleJ (see section 7.5, 7.6). Users can use them.

More specialized wrappers are also prepared for some cases.

The next wrapper is for the case that the multipole expansion is up to the monopole and the expansion center is the barycenter of particles.

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::MonopoleWithScatterSearch system;
```

The next wrapper is for the case that the multipole expansion is up to the quadrupole and the expansion center is the barycenter of particles.

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::QuadrupoleWithScatterSearch system;
```

The template arguments are the same as those for PS::SEARCH\_MODE\_LONG.

# 9.1.4.1.3 PS::SEARCH\_MODE\_LONG\_SYMMETRY

This section describes the wrapper class for PS::SEARCH\_MODE\_LONG\_SYMMETRY. Next example shows how to create an object of PS::SEARCH\_MODE\_LONG\_SYMMETRY class. Here, the object is named system.

```
PS::TreeForForceLong <TResult, TEpi, TEpj, TMom, TSpj>::WithSymmetrySearch system;
```

The template arguments are Force class, EssentialParticleI class, EssentialParticleJ class, Moment class for both local and global trees and SuperParticleJ class. FDPS prepare some useful classes of Moment and SuperParticleJ (see section 7.5, 7.6). Users can use them.

More specialized wrappers are also prepared for some cases.

The next wrapper is for the case that the multipole expansion is up to the monopole and the expansion center is the barycenter of particles.

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::MonopoleWithSymmetrySearch system;
```

The next wrapper is for the case that the multipole expansion is up to the quadrupole and the expansion center is the barycenter of particles.

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::QuadrupoleWithSymmetrySearch system;
```

The template arguments are the same as those for PS::SEARCH\_MODE\_LONG.

# 9.1.4.1.4 PS::SEARCH\_MODE\_LONG\_CUTOFF

This section describes the wrapper class for PS::SEARCH\_MODE\_LONG\_CUTOFF. Next example shows how to create an object of PS::SEARCH\_MODE\_LONG\_CUTOFF class. Here, the object is named system.

```
PS::TreeForForceLong <TResult, TEpi, TEpj, TMom, TSpj>::WithCutoff system;
```

The template arguments are Force class, EssentialParticleI class, EssentialParticleJ class, Moment class for both local and global trees and SuperParticleJ class. FDPS prepare some useful classes of Moment and SuperParticleJ (see section 7.5, 7.6). Users can use them.

There is the more specialized wrapper for the case that the multipole expansion is up to the monopole and the expansion center is the barycenter of particles.

```
PS::TreeForForceLong <TResult, TEpi, TEpj>::MonopoleWithCutoff system;
```

# 9.1.4.1.5 PS::SEARCH\_MODE\_GATHER

This wrapper is for the gather mode search. The template arguments are Force class, EssentialParticleI class and EssentialParticleJ class.

Next example shows how to create an object of PS::SEARCH\_MODE\_GATHER class. Here, the object is named system.

```
PS::TreeForForceShort<TResult, TEpi, TEpj>::Gather system;
```

# 9.1.4.1.6 PS::SEARCH\_MODE\_SCATTER

This wrapper is for the scatter mode search. The template arguments are Force class, EssentialParticleI class and EssentialParticleJ class.

Next example shows how to create an object of PS::SEARCH\_MODE\_SCATTER class. Here, the object is named system.

```
PS::TreeForForceShort<TResult, TEpi, TEpj>::Scatter system;
```

# 9.1.4.1.7 PS::SEARCH\_MODE\_SYMMETRY

This wrapper is for the symmetry mode search. The template arguments are Force class, EssentialParticleI class and EssentialParticleJ class.

Next example shows how to create an object of PS::SEARCH\_MODE\_SYMMETRY class. Here, the object is named system.

```
PS::TreeForForceShort<TResult, TEpi, TEpj>::Symmetry system;
```

### 9.1.4.2 API

This module has APIs for initial setup, calculation of interaction and searching neighbors. The following subsections describe them.

Next examples are declarations of member functions of the TreeForForce class.

```
template <class TSearchMode,
          class TResult,
          class TEpi,
          class TEpj,
          class TMomLocal,
          class TMomGlobal,
          class TSpj>
void PS::TreeForForce<TSearchMode,</pre>
                       TEpi,
                       TEpj,
                       TMomLocal,
                       TMomGlobal,
                       TSpj>::MemberFunction1();
template <class TSearchMode,
          class TResult,
          class TEpi,
          class TEpj,
          class TMomLocal,
          class TMomGlobal,
          class TSpj>
template <class TTT>
void PS::TreeForForce<TSearchMode,</pre>
                       TEpi,
                       TEpj,
                       TMomLocal,
                       TMomGlobal,
                       TSpj>::MemberFunction2(TTT arg1);
```

In this section, to simplify notation, the template arguments of the TreeForForce class are omitted as follows.

```
void PS::TreeForForce::MemberFunction1();

template <class TTT>
void PS::TreeForForce::MemberFunction2(TTT arg1);
```

# 9.1.4.2.1 Initial Setup

The APIs of initial setup of TreeForForce class are declared below.

# Listing 37: TreeForForce1

```
namespace ParticleSimulator {
1
2
       template < class TSearchMode,
3
                 class TResult,
4
                 class TEpi,
5
                 class TEpj,
6
                 class TMomLocal,
7
                 class TMomGlobal,
8
                 class TSpj>
9
       class TreeForForce{
10
       public:
11
       void TreeForForce();
12
       void initialize(const U64 n_glb_tot,
13
                         const F32 theta=0.7,
                         const U32 n_leaf_limit=8,
14
15
                         const U32 n_group_limit=64);
16
       };
17 }
```

# 9.1.4.2.1.1 Constructor

Constructor

```
void PS::TreeForForce();
```

• arguments

void.

• returned value

void.

• function

Create an object of TreeForForce class.

# 9.1.4.2.1.2 PS::TreeForForce::initialize

PS::TreeForForce::initialize

# • arguments

n\_glb\_tot: input. Type const PS::U64. Total number of particles for all processes.

theta: input. Type const PS::F32. Opening criterion of tree.

n\_leaf\_limit: input. Type const PS::U32. The maximum number of particles in leaf cell.

n\_group\_limit: input. Type const PS::U32. The maximum number of particles which share the same interactions list.

### • returned value

void.

### • function

Initialize an object of PS::TreeForForce class.

# 9.1.4.2.2 Low Level APIs

The low-level APIs for calculating forces are declared below.

# Listing 38: TreeForForce1

```
1 namespace ParticleSimulator {
2
       template < class TSearchMode,
3
                 class TResult,
4
                 class TEpi,
5
                 class TEpj,
6
                 class TMomLocal,
7
                 class TMomGlobal,
8
                 class TSpj>
       class TreeForForce{
9
10
       public:
11
           template < class Tpsys >
12
           void setParticleLocalTree(const Tpsys & psys,
13
                                        const bool clear=true);
14
           template < class Tdinfo >
15
           void makeLocalTree(const Tdinfo & dinfo);
           void makeLocalTree(const F32 1,
16
17
                                const F32vec & c = F32vec(0.0));
18
           template < class Tdinfo >
           void makeGlobalTree(const Tdinfo & dinfo);
19
20
           void calcMomentGlobalTree();
21
           template < class Tfunc_ep_ep >
22
           void calcForce(Tfunc_ep_ep pfunc_ep_ep,
                            const bool clear=true);
23
24
           template < class Tfunc_ep_ep, class Tfunc_sp_ep>
25
           void calcForce(Tfunc_ep_ep pfunc_ep_ep,
26
                            Tfunc_sp_ep pfunc_sp_ep,
```

```
const bool clear=true);
force getForce(const S32 i);
};
};
```

# 9.1.4.2.2.1 PS::TreeForForce::setParticleLocalTree

PS::TreeForForce::setParticleLocalTree

# • arguments

psys: input. Type const ParticleSystem. Object of ParticleSystem class clear: input. Type const bool. A flag to clear the data of particles, if set to true.

### • returned value

void.

# • function

Read the data of particles from psys and write them to the calling object. If clear is true, the data recorded are cleared.

# 9.1.4.2.2.2 PS::TreeForForce::makeLocalTree

PS::TreeForForce::makeLocalTree

# • arguments

dinfo: input. Type const DomainInfo &. Object of DomainInfo class.

## • returned value

void.

### • function

Make local tree. The root cell of the local tree is calculated by using dinfo.

# • arguments

1: input. Type const PS::F32. Length of a side of root cell of local tree.c: input. Type PS::F32vec. Center coordinate of root cell of local tree.

### • returned value

void.

### function

Make local tree. The two arguments must be the same for all processes. Otherwise, the results can be wrong.

# $9.1.4.2.2.3 \quad PS:: Tree For Force:: make Global Tree$

PS::TreeForForce::makeGlobalTree

# • arguments

dinfo: input. Type const DomainInfo &. An object of DomainInfo class.

# • returned value

void.

## • function

Make global tree.

# 9.1.4.2.2.4 PS::TreeForForce::calcMomentGlobalTree

PS::TreeForForce::calcMomentGlobalTree

```
void PS::TreeForForce::calcMomentGlobalTree();
```

## • arguments

void.

### • returned value

void.

### • function

Calculate moments of cells of global tree.

## 9.1.4.2.2.5 PS::TreeForForce::calcForce

PS::TreeForForce::calcForce

# • arguments

pfunc\_ep\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const EssentialParticleJ \* type, const PS::S32 type and Result \* type.

clear: input. Type const bool. A flag to clear the data of particles, if set to true.

### • returned value

void.

### • function

Calculate forces of all particles of the calling object. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_GATHER, PS::SEARCH\_MODE\_SCATTER and PS::SEARCH\_MODE\_SYMMETRY.

### • arguments

pfunc\_ep\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const EssentialParticleJ \* type, const PS::S32 type and Result \* type.

pfunc\_sp\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentailParticleI \* type, const PS::S32 type, const SuperParticleJ \* type, const PS::S32 type and Result \* type.

clear: input. Type const bool. A flag to clear the data of forces, if set to true.

### • returned value

void.

### function

Calculate forces of all particles loaded in the calling object. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_GATHER, PS::SEARCH\_MODE\_SCATTER or PS::SEARCH\_MODE\_SYMMETRY.

# 9.1.4.2.2.6 PS::TreeForForce::getForce

PS:: Tree For Force:: get Force

```
TResult PS::TreeForForce::getForce(const PS::S32 i);
```

## • arguments

i: input. Type const PS::S32. Index of array of particles.

### • returned value

Type Force. Force on the i-th particle.

### • function

Return the force on the i-th particle.

# 9.1.4.2.3 High Level APIs

The high level APIs for interactions are declared below.

# Listing 39: TreeForForce1

```
1 namespace ParticleSimulator {
2
       template < class TSearchMode,
3
                 class TResult,
4
                 class TEpi,
5
                 class TEpj,
6
                 class TMomLocal,
7
                 class TMomGlobal,
8
                 class TSpj>
       class TreeForForce{
9
10
       public:
11
            template < class Tfunc_ep_ep,
12
                      class Tpsys,
                      class Tdinfo>
13
            void calcForceAllAndWriteBack
14
15
                         (Tfunc_ep_ep pfunc_ep_ep,
16
                          Tpsys & psys,
17
                          Tdinfo & dinfo,
```

```
18
                         const bool clear_force = true,
                          const INTERACTION_LIST_MODE list_mode =
19
                               MAKE_LIST);
20
21
           template < class Tfunc_ep_ep,
22
                     class Tfunc_sp_ep,
23
                     class Tpsys,
                     class Tdinfo>
24
25
           void calcForceAllAndWriteBack
26
                        (Tfunc_ep_ep pfunc_ep_ep,
27
                         Tfunc_sp_ep pfunc_sp_ep,
28
                         Tpsys & psys,
29
                         TDinfo & dinfo,
                         const bool clear_force=true,
30
31
                         const INTERACTION_LIST_MODE list_mode =
                               MAKE_LIST);
32
33
34
           template < class Tfunc_dispatch,
35
                     class Tfunc_retrieve,
36
                     class Tpsys,
37
                     class Tdinfo>
38
           void calcForceAllandWriteBackMultiWalk
39
                 (Tfunc_dispatch pfunc_dispatch,
40
                  Tfunc_retrieve pfunc_retrieve,
41
                  Tpsys & psys,
42
                  Tdinfo & dinfo,
43
                  const S32 nwalk,
44
                  const bool clear=true,
                  const INTERACTION_LIST_MODE list_mode = MAKE_LIST
45
                        );
46
47
           void calcForceAllandWriteBackMultiWalkIndex
48
                 (Tfunc_dispatch pfunc_dispatch,
49
                  Tfunc_retrieve pfunc_retrieve,
50
                  Tpsys & psys,
51
                  Tdinfo & dinfo,
52
                  const S32 nwalk,
53
                  const bool clear=true,
                  const INTERACTION_LIST_MODE list_mode = MAKE_LIST
54
                        );
55
56
           template < class Tfunc_ep_ep,
57
                     class Tfunc_sp_ep,
58
                     class Tpsys,
59
                     class Tdinfo>
```

```
60
            void calcForceAll
61
                         (Tfunc_ep_ep pfunc_ep_ep,
62
                          Tfunc_sp_ep pfunc_sp_ep,
63
                          Tpsys & psys,
64
                          Tdinfo & dinfo,
65
                          const bool clear_force=true,
66
                          const INTERACTION_LIST_MODE list_mode =
                                MAKE_LIST);
67
68
            template < class Tfunc_ep_ep,
69
                      class Tdinfo>
70
            void calcForceAll(Tfunc_ep_ep pfunc_ep_ep,
71
                                Tfunc_sp_ep pfunc_sp_ep,
72
                                Tpsys & psys,
73
                                Tdinfo & dinfo,
74
                                const bool clear_force=true,
75
                                const INTERACTION_LIST_MODE list_mode
                                       = MAKE_LIST);
76
77
            template < class Tfunc_ep_ep,
78
                      class Tdinfo>
79
            void calcForceMakeingTree
80
                         (Tfunc_ep_ep pfunc_ep_ep,
81
                          Tdinfo & dinfo,
82
                          const bool clear_force=true,
83
                          const INTERACTION_LIST_MODE list_mode =
                                MAKE_LIST);
84
85
            template < class Tfunc_ep_ep,
                      class Tfunc_sp_ep,
86
87
                      class Tdinfo>
88
            void calcForceMakingTree
                         (Tfunc_ep_ep pfunc_ep_ep,
89
90
                          Tfunc_sp_ep pfunc_sp_ep,
91
                          Tdinfo & dinfo,
92
                          const bool clear_force=true,
93
                          const INTERACTION_LIST_MODE list_mode =
                                MAKE_LIST);
94
95
            template < class Tfunc_ep_ep,
96
                      class Tpsys>
97
            void calcForceAndWriteBack
98
                         (Tfunc_ep_ep pfunc_ep_ep,
                          Tpsys & psys,
99
100
                          const bool clear=true,
101
                          const INTERACTION_LIST_MODE list_mode =
```

```
MAKE_LIST);
102
103
            template < class Tfunc_ep_ep,
104
                      class Tfunc_sp_ep,
                      class Tpsys>
105
106
            void calcForceAndWriteBack
107
                          (Tfunc_ep_ep pfunc_ep_ep,
108
                           Tfunc_sp_ep pfunc_sp_ep,
109
                           Tpsys & psys,
110
                           const bool clear=true,
111
                           const INTERACTION_LIST_MODE list_mode =
                                MAKE_LIST);
112
        };
113 }
114 namespace PS = ParticleSimulator;
```

# 9.1.4.2.3.1 PS::TreeForForce::calcForceAllAndWriteBack PS::TreeForForce::calcForceAllAndWriteBack

#### • arguments

pfunc\_ep\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const EssentialParticleJ \* type, const PS::S32 type and Result \* type.

psys: input and output. Type ParticleSystem &. Object of ParticleSystem class.

dinfo: input. Type DomainInfo &. Object of DomainInfo class

clear: input. Type const bool. A flag to clear the data of forces if set to true.

list\_mode: input. Type const PS::INTERACTION\_LIST\_MODE. This argument controls the behavior of the API with respect to the reuse of interaction lists. The value must be

either of PS::MAKE\_LIST, PS::MAKE\_LIST\_FOR\_REUSE or PS::REUSE\_LIST. The action of the API is not determined for other values. If PS::MAKE\_LIST is given, FDPS makes interaction lists newly and performs interaction calculations using them. FDPS does not store these interaction lists. Hence, we cannot reuse these lists in the next interaction calculation (in the next call of the API). If PS::MAEK\_LIST\_FOR\_REUSE is given, FDPS makes interaction lists newly and stores them internally for future reuse. Then, FDPS performs interaction list calculation. Therefore, we can reuse the interaction lists in the next interaction calculation. When PS::REUSE\_LIST is given, FDPS performs interaction calculation using the interaction lists created previously with PS::MAEK\_LIST\_FOR\_REUSE. If list\_mode is omitted, FDPS acts as if PS::MAKE\_LIST is given.

#### • returned value

void.

#### • function

Calculate forces of all particles in psys and write back these forces to psys. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_GATHER, PS::SEARCH\_MODE\_SCATTER and PS::SEARCH\_MODE\_SYMMETRY.

By PS::INTERACTION\_LIST\_MODE, user program can reuse the same interaction list constructed before. During reusing the list, user program MUST NOT call PS::ParticleSystem::ex change order of particles or delete particles.

```
template < class Tfunc_ep_ep,
         class Tfunc_sp_ep,
         class Tpsys,
         class Tdinfo>
void PS::TreeForForce::calcForceAllandWriteBack
             (Tfunc_ep_ep pfunc_ep_ep(TEpi *,
                                       const PS::S32,
                                       TEpj *,
                                       const PS::S32,
                                       TResult *),
              Tfunc_sp_ep pfunc_sp_ep(TEpi *,
                                       const PS::S32,
                                       TSpj *,
                                       const PS::S32,
                                       TResult *),
              Tpsys & psys,
              Tdinfo & dinfo
              const bool clear=true,
              const INTERACTION_LIST_MODE list_mode = MAKE_LIST);
```

#### • arguments

pfunc\_ep\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const EssentialParticleJ \* type, const PS::S32 type and Result \* type.

pfunc\_sp\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const SuperParticleJ \* type, const PS::S32 type and Result \* type.

psys: input and output. Type ParticleSystem &. An object of ParticleSystem class.

dinfo: input. Type DomainInfo &. An object of DomainInfo class.

clear: input. Type const bool. A flag to clear the data of forces, if set to true.

list\_mode: input. Type const PS::INTERACTION\_LIST\_MODE. This argument controls the behavior of the API with respect to the reuse of interaction lists. The value must be either of PS::MAKE\_LIST, PS::MAKE\_LIST\_FOR\_REUSE or PS::REUSE\_LIST. The action of the API is not determined for other values. If PS::MAKE\_LIST is given, FDPS makes interaction lists newly and performs interaction calculations using them. FDPS does not store these interaction lists. Hence, we cannot reuse these lists in the next interaction calculation (in the next call of the API). If PS::MAEK\_LIST\_FOR\_REUSE is given, FDPS makes interaction lists newly and stores them internally for future reuse. Then, FDPS performs interaction list calculation. Therefore, we can reuse the interaction lists in the next interaction calculation. When PS::REUSE\_LIST is given, FDPS performs interaction calculation using the interaction lists created previously with PS::MAEK\_LIST\_FOR\_REUSE. If list\_mode is omitted, FDPS acts as if PS::MAKE\_LIST is given.

#### • arguments

void.

#### function

Calculate forces of particles in psys and write back these forces to psys. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_LONG or PS::SEARCH\_MODE\_LONG\_CUTOFF.

By PS::INTERACTION\_LIST\_MODE, user program can reuse the same interaction list constructed before. During reusing the list, user program MUST NOT call PS::ParticleSystem::ex change order of particles or delete particles.

## • Arguments

pfunc\_dispatch: Input. Returns void. A functor which receives the arrays of EssentialParticleI, EssentialParticleJ (and SuperParticleJ) and dispatch the interaction calculation. When PS::SEARCH\_MODE is PS::SEARCH\_MODE\_LONG or PS::SEARCH\_MODE\_LONG\_CUTOFF, this function should has the following form:

This function returns zero upon normal completion.

When PS::SEARCH\_MODE is PS::SEARCH\_MODE\_GATHER or PS::SEARCH\_MODE\_SCATTER, or PS::SEARCH\_MODE\_SYMMETRY, this function should has the following form:

```
PS::S32 pfunc_dispatch(const PS::S32 tag, const PS::S32 nwalk, const TEpi** iptcl, const PS::S32* ni, const TEpj** jptcl_ep, const PS::S32* nj_ep);
```

This function returns zero upon normal completion.

pfunc\_retrieve: Input. A functor of type void. Returns the result calculated by the call to pfunc\_dispatch. This function should have the following form:

Here, tag connects the call to pfunc\_dispatch() and that to pfunc\_retrieve(). The result of calculation by a call to pfunc\_dispatch is retrieved by the call to pfunc\_retrieve() with the same value of tag.

tag\_max: Input. Type const PS::S32. The maximum number of tags. Tag values between zero and tag\_max -1 are allowed. Non-positive value causes error. The current version ignores the value greater than unity and uses tag vale of zero only.

psys: input. Type Tpsys & An object of ParticleSystem class.

dinfo: input. Type DomainInfo &. An object of DomainInfo class.

nwalk: input. Type const PS::S32 The maximum number of interaction list to be sent by a single call to dispatch/retrive.

clear: inpuyt. Type const bool. If true, the interaction result array is cleared before starting the calculation. Default is true.

list\_mode: input. Type const PS::INTERACTION\_LIST\_MODE. This argument controls the behavior of the API with respect to the reuse of interaction lists. The value must be either of PS::MAKE\_LIST, PS::MAKE\_LIST\_FOR\_REUSE or PS::REUSE\_LIST. The action of the API is not determined for other values. If PS::MAKE\_LIST is given, FDPS makes interaction lists newly and performs interaction calculations using them. FDPS does not store these interaction lists. Hence, we cannot reuse these lists in the next interaction calculation (in the next call of the API). If PS::MAEK\_LIST\_FOR\_REUSE is given, FDPS makes interaction lists newly and stores them internally for future reuse. Then, FDPS performs interaction list calculation. Therefore, we can reuse the interaction lists in the next interaction calculation. When PS::REUSE\_LIST is given, FDPS performs interaction calculation using the interaction lists created previously with PS::MAEK\_LIST\_FOR\_REUSE. If list\_mode is omitted, FDPS acts as if PS::MAKE\_LIST is given.

#### • Return value

None

#### • Function

Calculate all interactions between all particles in psys and store the result to psys. The multiwalk method is used for the interaction calculation. The maximum number of the interaction list is nwalk.

By PS::INTERACTION\_LIST\_MODE, user program can reuse the same interaction list constructed before. During reusing the list, user program MUST NOT call PS::ParticleSystem::ex change order of particles or delete particles.

#### 9.1.4.2.3.2 PS::TreeForForce::calcForceAll

PS::TreeForForce::calcForceAll

#### • arguments

pfunc\_ep\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const EssentialParticleJ \* type, const PS::S32 type and Result \* type.

psys: input and output. Type ParticleSystem &. An object of ParticleSystem class.

dinfo: input. Type DomainInfo &. An object of DomainInfo class.

clear: input. Type const bool. A flag to clear the data of forces, if set to true.

list\_mode: input. Type const PS::INTERACTION\_LIST\_MODE. This argument controls the behavior of the API with respect to the reuse of interaction lists. The value must be either of PS::MAKE\_LIST, PS::MAKE\_LIST\_FOR\_REUSE or PS::REUSE\_LIST. The action of the API is not determined for other values. If PS::MAKE\_LIST is given, FDPS makes interaction lists newly and performs interaction calculations using them. FDPS does not store these interaction lists. Hence, we cannot reuse these lists in the next interaction calculation (in the next call of the API). If PS::MAEK\_LIST\_FOR\_REUSE is given, FDPS makes interaction lists newly and stores them internally for future reuse. Then, FDPS performs interaction list calculation. Therefore, we can reuse the interaction lists in the next interaction calculation. When PS::REUSE\_LIST is given, FDPS performs interaction calculation using the interaction lists created previously with PS::MAEK\_LIST\_FOR\_REUSE. If list\_mode is omitted, FDPS acts as if PS::MAKE\_LIST is given.

#### • returned value

void.

#### • function

Calculate forces of particles in psys. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_SYMMETRY. This API works similar as that of PS::TreeForForce::calcForceAllAndWriteBack without writing back forces to psys.

By PS::INTERACTION\_LIST\_MODE, user program can reuse the same interaction list constructed before. During reusing the list, user program MUST NOT call PS::ParticleSystem::ex change order of particles or delete particles.

```
template < class Tfunc_ep_ep,
         class Tfunc_sp_ep,
         class Tpsys,
         class Tdinfo>
void PS::TreeForForce::calcForceAll
             (Tfunc_ep_ep pfunc_ep_ep(TEpi *,
                                       const PS::S32,
                                       TEpj *,
                                       const PS::S32,
                                       TResult *),
              Tfunc_sp_ep pfunc_sp_ep(TEpi *,
                                       const PS::S32,
                                       TSpj *,
                                       const PS::S32,
                                       TResult *),
              Tpsys & psys,
              Tdinfo & dinfo
              const bool clear=true,
              const INTERACTION_LIST_MODE list_mode = MAKE_LIST);
```

#### arguments

pfunc\_ep\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const EssentialParticleJ \* type, const PS::S32 type and Result \* type.

pfunc\_sp\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const SuperParticleJ \* type, const PS::S32 type and Result \* type.

psys: input. Type ParticleSystem &. An object of ParticleSystem class

dinfo: input. Type DomainInfo &. An object of DomainInfo class

clear: input. Type const bool. A flag to clear the data of forces, if set to true.

list\_mode: input. Type const PS::INTERACTION\_LIST\_MODE. This argument controls the behavior of the API with respect to the reuse of interaction lists. The value must be either of PS::MAKE\_LIST, PS::MAKE\_LIST\_FOR\_REUSE or PS::REUSE\_LIST. The action of the API is not determined for other values. If PS::MAKE\_LIST is given, FDPS

makes interaction lists newly and performs interaction calculations using them. FDPS does not store these interaction lists. Hence, we cannot reuse these lists in the next interaction calculation (in the next call of the API). If PS::MAEK\_LIST\_FOR\_REUSE is given, FDPS makes interaction lists newly and stores them internally for future reuse. Then, FDPS performs interaction list calculation. Therefore, we can reuse the interaction lists in the next interaction calculation. When PS::REUSE\_LIST is given, FDPS performs interaction calculation using the interaction lists created previously with PS::MAEK\_LIST\_FOR\_REUSE. If list\_mode is omitted, FDPS acts as if PS::MAKE\_LIST is given.

#### • returned value

void.

# • function

Calculate forces of particles in psys. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_LONG\_CUTOFF. This API works similar as that of PS::TreeForForce::calcForceAllAndWriteBack without writing back forces to psys.

By PS::INTERACTION\_LIST\_MODE, user program can reuse the same interaction list constructed before. During reusing the list, user program MUST NOT call PS::ParticleSystem::ex change order of particles or delete particles.

# 9.1.4.2.3.3 PS::TreeForForce::calcForceAndWriteBack PS::TreeForForce::calcForceAndWriteBack

#### • arguments

pfunc\_ep\_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI \* type, const PS::S32 type, const EssentialParticleJ \* type, const PS::S32 type and Result \* type.

psys: input. Type ParticleSystem &.

clear: input. Type const bool. A flag to clear the data of forces, if set to true.

#### • returned value

void.

#### • function

Calculate forces of particles in psys by using the global tree created before. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_GATHER, PS::SEARCH\_MODE\_SCATTER or PS::SEARCH\_MODE\_SYMMETRY. This API works similar as that of PS::TreeForForce::calcForceAl without reading particle from psys, making the local tree, making global tree and calculate momentum.

```
template < class Tfunc_ep_ep,
         class Tfunc_sp_ep,
         class Tpsys>
void PS::TreeForForce::calcForceAllandWriteBack
             (Tfunc_ep_ep pfunc_ep_ep(TEpi *,
                                       const PS::S32,
                                        TEpj *,
                                        const PS::S32,
                                        TResult *),
              Tfunc_sp_ep pfunc_sp_ep(TEpi *,
                                        const PS::S32,
                                        TSpj *,
                                        const PS::S32,
                                       TResult *),
              Tpsys & psys,
              const bool clear=true);
```

#### • arguments

```
pfunc_ep_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI * type, const PS::S32 type, const EssentialParticleJ * type, const PS::S32 type and Result * type.

pfunc_sp_ep: input. A Functor of which a returned value is void and of which arguments are const EssentialParticleI * type, const PS::S32 type, const SuperParticleJ * type, const PS::S32 type and Result * type.

psys: output. Type ParticleSystem &.

clear: input. Type const bool. A flag to clear the data of forces, if set to true.
```

# • returned value

void.

#### function

Calculate forces of particles in psys by using the global tree created previously. This function can be used only if PS::SEARCH\_MODE is PS::SEARCH\_MODE\_LONG\_C

This API works similar as that of PS::TreeForForce::calcForceAllAndWriteBack without reading particle from psys, making the local tree, making global tree and calculate momentum.

9.1.4.2.4 Neighbor List

9.1.4.2.4.1 getNeighborListOneParticle getNeighborListOneParticle

```
template<class Tptcl>
PS::S32 PS::TreeForForce::getNeighborListOneParticle(const Tptcl & ptcl,
                                                      EPJ * \& epj);
```

#### arguments

ptcl: input. Type Tptcl &. A particle to obtain its neighbor particles.

epj: output. Type EssentailParticleJ \* &. The pointer to the beginning of the array of neighbor particles of ptcl.

#### returned value

Type PS::S32 &. The number of neighbor particles of ptcl.

#### • function

By using the global tree of the calling object, search the neighbors of ptcl and set the pointer to the beginning of the array of neighbors to epj and return the number of neighbors. Note that epj is the pointer to the first element of the array of the particle data of type EPJ (not the array of pointers to particles), and points to the internal buffer maintained by FDPS. Therefore, users must not call free() or delete() to epj. This function is thread safe. In other words, each thread has its own buffer. Since each thread has only one buffer, the content of the buffer is overwritten at each call to this function.

This function is available only if PS::SEARCH\_MODE is any one of

- PS::SEARCH\_MODE\_GATHER,
- PS::SEARCH\_MODE\_SCATTER,PS::SEARCH\_MODE\_SYMMETRY,
- PS::SEARCH\_MODE\_LONG\_SCATTER,
- PS::SEARCH\_MODE\_LONG\_SYMMETRY,
- PS::SEARCH\_MODE\_LONG\_GATHER (not implemented yet)
- PS::SEARCH\_MODE\_LONG\_CUTOFF\_GATHER (not implemented yet),
- PS::SEARCH\_MODE\_LONG\_CUTOFF\_SCATTER (not implemented yet),
- and PS::SEARCH\_MODE\_LONG\_CUTOFF\_SYMMETRY (not implemented yet).

ptcl needs a member function PS::F64vec getPos() like FullParticle class. If PS::SEARCH\_MODE is any one of

- PS::SEARCH\_MODE\_GATHER,
- PS::SEARCH\_MODE\_SYMMETRY,

```
PS::SEARCH_MODE_LONG_SYMMETRY,
PS::SEARCH_MODE_LONG_GATHER (not implemented yet),
PS::SEARCH_MODE_LONG_CUTOFF_GATHER (not implemented yet),
and PS::SEARCH_MODE_LONG_CUTOFF_SYMMETRY (not implemented yet),
```

ptcl also needs a member function PS::F64 getRSearch().

# 9.1.4.2.5 Time Measurment

The APIs for time measurment are declared below. When a member function is called, its execution time is set to the private member time\_profile. Until the method PS::TreeForForce::clearTimeProfile() is called, execution times are accumulated.

# Listing 40: TreeForForce2

```
1
  namespace ParticleSimulator {
2
       template < class TSearchMode,
3
                 class TResult,
4
                 class TEpi,
5
                 class TEpj,
6
                 class TMomLocal,
7
                 class TMomGlobal,
8
                 class TSpj>
9
       class TreeForForce{
10
       public:
            TimeProfile getTimeProfile();
11
            void clearTimeProfile();
12
13
       };
14 }
```

## 9.1.4.2.5.1 PS::TreeForForce::getTimeProfile

PS::TreeForForce::getTimeProfile

```
PS::TimeProfile PS::TreeForForce::getTimeProfile();
```

#### • arguments

void.

#### • returned value

Type PS::TimeProfile.

# • function

The execution time for creation of local tree, creation of global tree, evaluation of force, evaluation of momenta of local tree, evaluation of momenta of global tree, creation of LET and exchange LET are recorded appropriate members of the private member variable time\_profile\_, make\_local\_tree, make\_global\_tree\_, calc\_force\_,

calc\_moment\_local\_tree\_, calc\_moment\_global\_tree\_, make\_LET\_1st\_, make\_LET\_2nd\_,
exchange\_LET\_1st\_ and exchange\_LET\_2nd\_.

# 9.1.4.2.5.2 PS::TreeForForce::clearTimeProfile

PS::TreeForForce::clearTimeProfile

```
void PS::TreeForForce::clearTimeProfile();
```

• arguments

void.

• returned value

void.

• function

Set all member variables of time\_profile\_ to 0.

#### 9.1.4.2.6 Obtain Information

The member functions of obtaining information are declared below.

Listing 41: TreeForForce2

```
1 namespace ParticleSimulator {
2
       template < class TSearchMode,
3
                 class TResult,
4
                 class TEpi,
5
                 class TEpj,
6
                 class TMomLocal,
7
                 class TMomGlobal,
8
                 class TSpj>
9
       class TreeForForce{
10
       public:
11
           Count_t getNumberOfInteractionEPEPLocal();
12
           Count_t getNumberOfInteractionEPSPLocal();
           Count_t getNumberOfInteractionEPEPGlobal();
13
14
           Count_t getNumberOfInteractionEPSPGlobal();
           void clearNumberOfInteraction();
15
16
           S64 getMemSizeTotalUsed();
17
       };
18
```

#### 9.1.4.2.6.1 PS::TreeForForce::getNumberOfInteractionEPEPLocal

PS:: Tree For Force:: get Number Of Interaction EPEP Local

PS::CountT PS::TreeForForce::getNumberOfInteractionEPEPLocal();

# • arguments

void.

#### • returned value

Type PS::CountT.

#### • function

Return the number of interactions between EssentialParticleI and EssentialParticleJ in the calling process.

# $9.1.4.2.6.2 \quad PS:: Tree For Force:: get Number Of Interaction EPEP Global$

PS:: Tree For Force:: get Number Of Interaction EPEPG lobal

PS::CountT PS::TreeForForce::getNumberOfInteractionEPEPGlobal();

# • arguments

void.

# • returned value

Type PS::CountT.

#### • function

Return the total number of interactions between  ${\tt EssentialParticleI}$  and  ${\tt EssentialParticleJ}$ , evaluated in all processes.

# $9.1.4.2.6.3 \quad PS:: TreeForForce:: getNumberOfInteraction EPSPL ocal$

PS::TreeForForce::getNumberOfInteractionEPSPLocal

PS::Count\\_t PS::TreeForForce::getNumberOfInteractionEPSPLocal();

#### • arguments

void.

# • returned value

Type PS::CountT.

#### • function

Return the number of interactions between EssentialParticleI and SuperParticleJ in the calling process.

# $9.1.4.2.6.4 \quad PS:: Tree For Force:: get Number Of Interaction EPSP Global$

PS:: Tree For Force:: get Number Of Interaction EPSP Global

PS::S64 PS::TreeForForce::getNumberOfInteractionEPSPGlobal();

# • arguments

void.

#### • returned value

Type PS::CountT.

#### • function

Return the total number of interactions between EssentialParticleI and SPJ, evaluated in all processes.

# 9.1.4.2.6.5 PS::TreeForForce::clearNumberOfInteraction

PS:: Tree For Force:: clear Number Of Interaction

void PS::TreeForForce::clearNumberOfInteraction();

# • arguments

void.

#### • returned value

void.

#### • function

The counter for recording the number of interactions is set to 0.

# 9.1.4.2.6.6 PS::TreeForForce::getNumberOfWalkLocal

PS:: Tree For Force:: get Number Of Walk Local

PS::Count\\_t PS::TreeForForce::getNumberOfWalkLocal();

#### • arguments

void.

#### • returned value

Type PS::CountT.

#### • function

Return the number of tree traverse for the calling process.

# $9.1.4.2.6.7 \quad PS:: TreeForForce:: getNumberOfWalkGlobal$

PS:: Tree For Force:: get Number Of Walk Global

PS::Count\\_t PS::TreeForForce::getNumberOfWalkGlobal();

# • arguments

void.

#### • returned value

Type PS::S64.

#### • function

Return the total number of tree traverse for all processes.

## 9.1.4.2.6.8 PS::TreeForForce::getMemSizeUsed

PS::TreeForForce::getMemSizeUsed

PS::S64 PS::TreeForForce::getMemSizeUsed();

#### • arguments

void.

#### • returned value

Type PS::S64.

#### • function

Return the size of used memory of an object in the unit of byte.

#### 9.1.4.2.7 Obtain EPJ from particle id

# $\begin{array}{ccc} 9.1.4.2.7.1 & getEpjFromId\\ & \text{getEpjFromId} \end{array}$

```
EPJ * PS::TreeForForce::getEpjFromId(const PS::S64 id)
```

# • arguments

id: input. Type const PS::S64. Index of Particle which user program want to obtain (The value returned by a member function getId() of EPJ).

#### • returned value

Type EPJ\*: Pointer of EPJ corresponding to id.

#### • function

This API is usable *only when* EPJ type has a member function getId(). This API returns the pointer to a EPJ whose particle ID is id. If id is not in the list of EPJ, the API just returns NULL. The action of the API is not determined for the case that EPJ more than one have the same ID. For more information of getId(), please see section 7.4.4.4.1.

#### 9.1.5 Comm Class

This section describes PS::Comm class which is a module used to communicate data among processes and keep the data for communication. Since the singleton pattern is applied in this class, users do not need to create an object.

#### 9.1.5.1 API

The APIs are declared below.

Listing 42: Communication

```
namespace ParticleSimulator {
1
2
       class Comm{
3
       public:
4
           static S32 getRank();
5
           static S32 getNumberOfProc();
           static S32 getRankMultiDim(const S32 id);
6
7
           static S32 getNumberOfProcMultiDim(const S32 id);
8
           static bool synchronizeConditionalBranchAND
9
                        (const bool local);
10
           static bool synchronizeConditionalBranchOR
11
                        (const bool local);
12
           template < class T>
13
           static T getMinValue(const T val);
           template < class Tfloat, class Tint>
14
           static void getMinValue(const Tfloat f_in,
15
16
                                      const Tint i_in,
17
                                     Tfloat & f_out,
                                     Tint & i_out);
18
19
           template < class T>
20
           static T getMaxValue(const T val);
21
           template < class Tfloat, class Tint>
22
           static void getMaxValue(const Tfloat f_in,
23
                                      const Tint i_in,
24
                                     Tfloat & f_out,
                                     Tint & i_out );
25
26
           template < class T>
27
           static T getSum(const T val);
28
           template < class T>
```

#### 9.1.5.1.1 *PS::Comm::getRank*

```
static PS::S32 PS::Comm::getRank();
```

• arguments

void.

• returned value

Type PS::S32.

• function

Return the rank of the calling process.

# 9.1.5.1.2 PS::Comm::getNumberOfProc

```
static PS::S32 PS::Comm::getNumberOfProc();
```

 $\bullet$  arguments

void.

• returned value

Type PS::S32. The total number of processes.

• function

Return the total number of processes.

# 9.1.5.1.3 PS::Comm::getRankMultiDim

```
static PS::S32 PS::Comm::getRankMultiDim(const PS::S32 id);
```

• arguments

id: input. Type const PS::S32. Id of axes. x-axis:0, y-axis:1, z-axis:2.

• returned value

Type PS::S32. The rank of the calling process along id-th axis. In the case of two dimensional simulations, FDPS returns 0 for id=2.

# 9.1.5.1.4 PS::Comm::getNumberOfProcMultiDim

static PS::S32 PS::Comm::getNumberOfProcMultiDim(const PS::S32 id);

• id: input. Type const PS::S32. id of axes. x-axis:0, y-axis:1, z-axis:2.

#### • returned value

Type PS::S32. The number of processes along id-th axis. In the case of two dimensional simulations, FDPS returns 1 for id=2.

## 9.1.5.1.5 PS::Comm::synchronizeConditionalBranchAND

static bool PS::Comm::synchronizeConditionalBranchAND(const bool local)

#### • arguments

local: input. Type const bool.

# • returned value

bool type. Logical product of local over all processes.

#### • function

Return logical product of local over all processes.

#### 9.1.5.1.6 PS::Comm::synchronizeConditionalBranchOR

static bool PS::Comm::synchronizeConditionalBranchOR(const bool local);

#### • arguments

local: input. Type const bool.

#### • returned value

Type bool.

#### • function

Return logical sum of local over all processes.

# 9.1.5.1.7 PS::Comm::getMinValue

```
template <class T>
static T PS::Comm::getMinValue(const T val);
```

#### • arguments

```
val: input. Type const T. Type T is allowed to be PS::F32, PS::F64, PS::S32, PS::S64, PS::U32 and PS::U64
```

#### • returned value

Type T. The minimum value of val of all processes.

#### • arguments

```
f_in: input. Type const Tfloat.
```

i\_in: input. Type const Tint.

f\_out: output. Type Tfloat &. Tfloat must be PS::F64 or PS::F32. The minimum value of f\_in among all processes.

i\_out: output. Type Tint &. Tint must be PS::S64, PS::S32, PS::U64 or PS::U32. The rank of the process of which f\_in is the minimum.

#### • returned value

void.

#### 9.1.5.1.8 PS::Comm::getMaxValue

```
template <class T>
static T PS::Comm::getMaxValue(const T val);
```

#### • arguments

val: input. Type const T.

#### • returned value

Type T. The maximum value of val of all processes.

## • arguments

f\_in: input. Type const Tfloat. Type Tfloat is PS::F32 or PS::F64.
i\_in: input. Type const Tint. Type Tint is PS::S32.
f\_out: output. Type Tfloat &. The maximum value of f\_in among all processes.

i\_out: output. Type Tint &. The rank of the process of which f\_in is the maximum.

#### • returned value

void.

# 9.1.5.1.9 PS::Comm::getSum

```
template <class T>
static T PS::Comm::getSum(const T val);
```

#### • arguments

val: input. Type const T. Type T is allowed to be PS::S32, PS::S64, PS::F32,
PS::F64, PS::U32, PS::U64, PS::F32vec and F64vec.

## • returned value

Type T. Return the global sum of val.

#### 9.1.5.1.10 PS::Comm::broadcast

#### • arguments

val: input. Type T \*. Type T is allowed to be any types.n: input. Type const PS::S32. The number of variables.src: input. Type const PS::S32. The rank of source process.

#### • returned value

void.

#### • function

Broadcast val for the src-th process.

#### 9.1.6 Other functions

In this section, functions defined directly in the namespace ParticleSimulator are described.

#### 9.1.6.1 Timing function

#### 9.1.6.1.1 PS::GetWtime

```
inline PS::F64 PS::GetWtime();
```

#### • Arguments

None.

#### • Return value

Type PS::F64. Wall-clock time in seconds.

#### 9.2 Extended Classes

# 9.2.1 Summary

This section describes two extended modules: one is the class for the Particle Mesh scheme and the other is the class for Phantom-GRAPE.

#### 9.2.2 Particle Mesh Class

This section describes ParticleMesh class which is for the evaluation of interactions between particles using the Particle Mesh method. The cutoff of the S-2 type function is applied to the mesh force, and the cutoff radius is fixed to three times the mesh size. The coordinates of particles sent to ParticleMesh class should be in the range of [0,1). The following subsections describe how to create the object for Particle Mesh method, APIs, predefined macros and how to use the class.

# 9.2.2.1 Creation of Object

ParticleMesh class is declared bellow.

```
Listing 43: ParticleMesh0
```

```
1 namespace ParticleSimulator {
2     namespace ParticleMesh {
```

```
egin{array}{lll} 3 & & & & & & & \\ 4 & & & & & \\ 5 & & & & & \\ \end{array}
```

Next example shows how to create an object of ParticleMesh class. Here, the object is named pm.

```
PS::PM::ParticleMesh pm;
```

#### 9.2.2.2 API

ParticleMesh class has APIs for initialization and force evaluation. The following subsections describe them.

#### 9.2.2.2.1 Initial Setup

# Listing 44: ParticleMesh1

```
1 namespace ParticleSimulator {
2    namespace ParticleMesh {
3         class ParticleMesh{
4             ParticleMesh();
5         };
6    }
7 }
```

#### 9.2.2.2.1.1 Constructor

Constructor

```
void PS::PM::ParticleMesh::ParticleMesh();
```

• arguments

void.

• returned value

void.

• function

Create an object of ParticleMesh class.

#### 9.2.2.2.2 Low Level API

The low Level APIs are declared below.

# Listing 45: ParticleMesh1

```
namespace ParticleSimulator {
1
2
       namespace ParticleMesh {
3
           class ParticleMesh{
4
                template < class Tdinfo >
5
                void setDomainInfoParticleMesh
6
                             (const Tdinfo & dinfo);
7
                template < class Tpsys >
8
                void setParticleParticleMesh
9
                             (const Tpsys & psys,
10
                              const bool clear=true);
11
                void calcMeshForceOnly();
12
                F32vec getForce(F32vec pos);
13
                F32 getPotential(F32vec pos);
14
           };
       }
15
16 }
```

9.2.2.2.2.1 PS::PM::ParticleMesh::setDomainInfoParticleMesh
PS::PM::ParticleMesh::setDomainInfoParticleMesh

• arguments

dinfo: input. Type const DomainInfo &. An object of DomainInfo class.

• returned value

void.

• function

Read the data from dinfo.

 $9.2.2.2.2.2 \quad PS::PM::ParticleMesh::setParticleParticleMesh$ 

PS::PM::ParticleMesh::setParticleParticleMesh

### • arguments

psys: input. Type const ParticleSystem &. An object of ParticleSystem class. clear: input. Type const bool. A flag to determine if previous information of particles loaded is cleared.

#### • returned value

void.

#### • function

Read data of particles from psys.

# $9.2.2.2.2.3 \quad PS::PM::ParticleMesh::calcMeshForceOnly$

PS::PM::ParticleMesh::calcMeshForceOnly

```
void PS::PM::ParticleMesh::calcMeshForceOnly();
```

## • arguments

void.

#### • returned value

void.

#### • function

Calculate forces on grid points.

# 9.2.2.2.2.4 PS::PM::ParticleMesh::getForce

PS::PM::ParticleMesh::getForce

```
PS::F32vec PS::PM::ParticleMesh::getForce(PS::F32vec pos);
```

# • arguments

pos: input. Type PS::F32vec.

# • returned value

Type PS::F32vec. PM force at the position pos.

#### • function

Return a PM force exerted at the position "pos". This function is thread-safe.

# $9.2.2.2.2.5 \quad PS::PM::ParticleMesh::getPotential$

PS::PM::ParticleMesh::getPotential

```
PS::F32 PS::PM::ParticleMesh::getPotential(PS::F32vec pos);
```

## • arguments

pos: input. Type PS::F32vec.

#### • returned value

Type PS::F32vec. PM potential at the position pos.

#### • function

Return a PM potential exerted at the position "pos". This function is thread-safe.

# 9.2.2.2.3 High Level API

High level APIs are declared below.

# Listing 46: ParticleMesh1

```
namespace ParticleSimulator {
1
2
       namespace ParticleMesh {
3
           class ParticleMesh{
4
                template < class Tpsys,
5
                          class Tdinfo>
6
                void calcForceAllAndWriteBack
7
                             (Tpsys & psys,
8
                              const Tdinfo & dinfo);
9
           };
10
       }
11 }
```

#### 9.2.2.2.3.1 PS::PM::ParticleMesh::calcForceAllAndWriteBack

PS::PM::ParticleMesh::calcForceAllAndWriteBack

#### • arguments

psys: input and output. Type ParticleSystem &. An object of the ParticleSystem class.

dinfo: input. Type const DomainInfo &. An object of the DomainInfo class.

• returned value

void.

• function

Calculate forces between particles of psys and write back the forces to psys.

#### 9.2.2.3 Predefined Macros

There are many macros predefined in this module and users must not define the same name macros. The predefined macros are listed bellow in alphabetical order.

- BINARY\_BOUNDARY
- BOUNDARY\_COMM\_NONBLOCKING
- BOUNDARY\_SMOOTHING
- BUFFER\_FOR\_TREE
- CALCPOT
- CLEAN\_BOUNDARY\_PARTICLE
- CONSTANT\_TIMESTEP
- EXCHANGE\_COMM\_NONBLOCKING
- FFT3D
- FFTW3\_PARALLEL
- FFTW\_DOUBLE
- FIX\_FFTNODE
- GADGET\_IO
- GRAPE\_OFF
- KCOMPUTER
- LONG\_ID
- MAKE\_LIST\_PROF
- MERGE\_SNAPSHOT
- MULTI\_TIMESTEP
- MY\_MPI\_BARRIER

- N128\_2H
- N256\_2H
- N256\_H
- N32\_2H
- N512\_2H
- NEW\_DECOMPOSITION
- NOACC
- NPART\_DIFFERENT\_DUMP
- OMP\_SCHDULE\_DISABLE
- PRINT\_TANIKAWA
- REVERSE\_ENDIAN\_INPUT
- REVERSE\_ENDIAN\_OUTPUT
- RMM\_PM
- SHIFT\_INITIAL\_BOUNDARY
- STATIC\_ARRAY
- TREE2
- TREECONSTRUCTION\_PARALLEL
- TREE\_PARTICLE\_CACHE
- UNIFORM
- UNSTABLE
- USING\_MPI\_PARTICLE
- VERBOSE\_MODE
- VERBOSE\_MODE2。

#### 9.2.2.4 How To Use Particle Mesh Class

Through the following steps, one can use this ParticleMesh class.

- 1. Compile ParticleMesh class.
- 2. Write a program including FDPS.
- 3. Compile the program.

The following subsections describe these steps.

#### 9.2.2.4.1 Compile of Particle Mesh Class

Users need to edit the Makefile in \$(FDPS)/src/particle\_mesh appropriately as follows.

- Set the macro INCLUDE\_FFTW (the directory in which the header file of FFTW is) in Makefile.
- Set the macro SIZE\_OF\_MESH (the number of meshes along one direction) in param\_fdps.h. Recommended value is  $N^{1/3}/2$  (N is the number of the particle).

After editing, users run make command.

If successful, the library libpm.a and the header file particle\_mesh.hpp are created.

#### 9.2.2.4.2 Writing Source Code

Next, users write programs as follows.

- Include the header files created above.
- Add member functions described below to FullParticle class (here, this particle class called FP).
  - void FP::copyFromForceParticleMesh(const PS::F32vec & force). In this function, copy force to arbitrary member variables in FP.
  - PS::F64 FP::getChargeParticleMesh(). This function returns mass.

#### 9.2.2.4.3 Compile of Source Code

To compile the programs written above, users need to follow the following steps.

- Specify the path to the header file particle\_mesh.hpp.
- Link the library libpm.a.
- Specify the path to the directory in which the header files of FFTW are.
- Link the FFTW libraries.

# 9.2.2.4.4 Note

ParticleMesh class does not work if the total number of processes is one.

#### 9.2.3 Phantom-GRAPE for X86

To use the low precision Phantom-GRAPE, refer to Tanikawa et al. (2013, New Astronomy, 19, 74). In this library, interactions with cutoff is also available. To use the high precision Phantom-GRAPE, refer to Tanikawa et al. (2012, New Astronomy, 17, 82).

# 10 Detection of Errors

#### 10.1 Abstract

FDPS equips features to detect compile time and run time errors. We describe errors detectable by FDPS, and how to deal with these errors. Note that unknown errors possibly happen. At that time, please inform us.

# 10.2 Compile time errors

(Not written yet)

#### 10.3 Run time errors

In run time error, FDPS outputs error messages in the following format, and terminates the program by PS::Abort(-1).

PS\_ERROR: ERROR MESSAGE

function: FUNCTION NAME, line: LINE NUMBER, file: FILE NAME

• ERROR MESSAGE

Error message.

• FUNCTION NAME

Function name in which an error happens.

• LINE NUMBER

Line number in which an error happens.

• FILE NAME

File name in which an error happens.

We list run time errors below.

#### 10.3.1 PS\_ERROR: can not open input file

This message indicates that there is no input file specified by users, when the file input functions in FDPS are called.

The following message follows the error message.

input file: "input file name"

# 10.3.2 PS\_ERROR: can not open output file

This message indiates that there is no output file specified by users, when the file output functions in FDPS are called.

The following message follows the error message.

output file: "output file name"

#### 10.3.3 PS\_ERROR: Do not initialize the tree twice

This message indicates that PS::TreeForForce::initialize(...) is called twice for the same tree object. Users can call this function only once.

#### 10.3.4 PS\_ERROR: The opening criterion of the tree must be >=0.0

This message indicates that a negative value is input into opening criterion of tree for long-distance force modes.

The following message follows the above message to the standard error.

theta\_= "input value for opening criterion"

SEARCH\_MODE: "data type for search mode"

Force: "Type name for tree force"

EPI: "type name for EPI"

EPJ: "type name for EPJ"

SPJ: "type name for SPJ"

# 10.3.5 PS\_ERROR: The limit number of the particles in the leaf cell must be > 0

This message indicates that a negative value is input into maximum particle number for a tree leaf cell.

The following message follows the above message to the standard error.

n\_leaf\_limit\_="input value for the maximum particle number in tree leaf cell"

SEARCH\_MODE: "data type for search mode"

Force: "Type name for tree force"

EPI: "type name for EPI"

EPJ: "type name for EPJ"

SPJ: "type name for SPJ"

# 10.3.6 PS\_ERROR: The limit number of particles in ip groups msut be >= that in leaf cells

This message indicates that the maximum particle number for a leaf cell is more than the maximum particle number for a i-group particle number, and when long-distant force modes are chosen.

The following message follows the above message to the standard error.

n\_leaf\_limit\_="Input the maximum particle number in a leaf cell"

n\_grp\_limit\_="Input the maximum particle number in a *i*-group particle number"

SEARCH\_MODE: "data type for search mode"

Force: "Type name for tree force"

EPI: "type name for EPI" EPJ: "type name for EPJ" SPJ: "type name for SPJ"

# 10.3.7 PS\_ERROR: The number of particles of this process is beyond the FDPS limit number

This message indicates that users deal with more than  $2G(G=2^{30})$  particles per MPI process.

# 10.3.8 PS\_ERROR: The forces w/o cutoff can be evaluated only under the open boundary condition

Thie message indicates that users set long-distance force without cutoff under periodic boundary condition.

# 10.3.9 PS\_ERROR: A particle is out of root domain

This message indicates that when the user program set the root domain by using function PS::DomainInfo::setRootDomain(...), any particle is outside the root domain. Particularly under periodic boudnary condition, the user program should shift particles from outside of the root domain to inside. For this purpose, we recommend the use of function PS::ParticleSystem::adjustPositionIntoRootDomain(...).

The following message follows the above message to the standard error.

position of the particle="position of particles outside" position of the root domain="coordinates of the root domain"

# 10.3.10 PS\_ERROR: The smoothing factor of an exponential moving average is must between 0 and 1.

This message indicates that users set the value which is less than 0 or grater than 1 as the smoothing factor of an exponential moving average by using function

PS::DomainInfo::initialize(...).

The following message follows the above message to the standard error.

The smoothing factor of an exponential moving average="The smoothing factor"

#### 10.3.11 PS\_ERROR: The coodinate of the root domain is inconsistent.

This message indicates that users set the coordinate of the lower vertex to be greater than the corresponding coordinate of the higher vertex by using function

PS::DomainInfo::setPosRootDomain(...).

The following message follows the above message for the standard error.

The coordinate of the low vertex of the rood domain="The coordinate of the lower vertex" The coordinate of the high vertex of the rood domain="The coordinate of the higher vertex"

#### 10.3.12 PS\_ERROR: Vector invalid accesse

This message indicates that users refer to an invalid element in the Vector class by using the operator "[]".

The following message follows the above message for the standard error.

Vector element="element user reffered" is not valid

# 11 Known Bugs

# 12 Limitation

 $\bullet$  Safe performance of integer types unique to FDPS is ensured, only when users adopt GCC and K compilers.

# 13 User Support

# 14 License

This software is MIT licensed. Please cite Iwasawa et al. (2016, Publications of the Astronomical Society of Japan, 68, 54) and Namekata et al. (2018, Publications of the Astronomical Society of Japan, 70, 70) if you use the standard functions only.

The extended feature "Particle Mesh" is implemented by using a module of GREEM code (Developers: Tomoaki Ishiyama and Keigo Nitadori) (Ishiyama, Fukushige & Makino 2009, Publications of the Astronomical Society of Japan, 61, 1319; Ishiyama, Nitadori & Makino, 2012 SC'12 Proceedings of the International Conference on High Performance Computing, Networking Stroage and Analysis, No. 5). GREEM code is developed based on the code in Yoshikawa & Fukushige (2005, Publications of the Astronomical Society of Japan, 57, 849). Please cite these three literatures if you use the extended feature "Particle Mesh".

Please cite Tanikawa et al. (2012, New Astronomy, 17, 82) and Tanikawa et al. (2012, New Astronomy, 19, 74) if you use the extended feature "Phantom-GRAPE for x86".

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# 15 Change Log

- 2015.03.13
  - Correct getRsearch to getRSearch. (section 7).
  - Add a function PS::Abort (section 8).
- 2015.03.17
  - Release version 1.0
- 2015.03.18
  - Modify the licence related to class Particle Mesh.
- 2015.03.20
  - Add the description of PS::Comm::broadcast.
- 2015.04.01
  - Add a caution that class Particle Mesh does not work in the case of 1 MPI process.
- 2015.10.07
  - Add the description of PM method. Section 9.2.2.
- 2015.12.01
  - Add the description of Multiwalk method. Sections A.10, A.11, 9.1.4.2.3.1, 7.11,
     7.12.
- 2016.02.19
  - Added descriptions for getNeighborListOneParticle in 9.1.4.2.4.
  - In accordance with the above change, added the descriptions of PS:: MomentMonopoleScatter and PS::MomentQuadrupoleScatter in section A.4.2.2.
  - Added the descriptions of their wrappers, MonopoleWithScatter and QuadrupoleWithScatter, to section 9.1.4.1.1.
- 2016.09.13
  - Added APIs for removing or adding particles. The discription of them is added in 9.1.3.2.5.
  - The discription of out-of-range access in type Vector is added in section 10.3.12.
- 2016.10.11
  - The implementation of the operator "[]" in the Vector class is cannged. Added remarks of the Vector class in section 6.4.

#### 2016.11.04

- Added APIs for IO in sections 9.1.3.2.3.1,9.1.3.2.3.2,9.1.3.2.3.3,9.1.3.2.3.4.

#### • 2017.07.11

- Added a discription about reusing interaction list in section ??.

#### • 2017.09.06

- Added API for sorting FP in particle system. The discription of this API is in section 9.1.3.2.7.3.
- Added API for obtaining a pointer of EPJ from its id. The discription of this API is in section 7.4.4.4.1,9.1.4.2.7.

#### • 2017.11.08

- Release FDPS version 4.0.

#### • 2018.01.16

- Release FDPS version 4.0b.
  - \* Fix a bug for memory allocation.
  - \* Add destructors for TreeForForce class, etc.

#### • 2018.05.25

- Release FDPS version 4.0c.
  - \* MPI C++ bindings in Particle Mesh extension are eliminated.

#### • 2018.06.13

- Release FDPS version 4.1.
  - \* Fix an issue when  $\theta = 0$  is specified (FDPS makes superparticles in some situations).
  - \* Description of orthotope type is added in Section 6.5.
  - \* Description of required member functions for specific cases is added in Section of Moment class.

#### • 2018.06.20

- Release FDPS version 4.1a.
  - \* Fix an omission in this section.

#### • 2018.08.2

- A new API getBoundaryCondition is added.
- The meaning of the argument weight of API collectSampleParticle is described in detail.

# • 2018.12.7

- Release FDPS version 5.0a.
  - \* Descriptions on pre-defined classes for

PS::SEARCH\_MODE\_LONG\_SCATTER

PS::SEARCH\_MODE\_LONG\_SYMMETRY

are added to the sections of Moment and SuperParticleJ classes ( $\S$  7.5 and  $\S$  7.6).

\* Descriptions on member functions readBinary and writeBinary are added to the sections of FullParticle and header classes (§ 7.2 and § 7.8). Examples of implementations of them are also added to § A.1 and § A.7.

#### • 2018.12.9

- Release FDPS 5.0b.
  - \* In accurate description of API getNeighborListOneParticle is fixed (No changes in the source codes of FDPS except for the version information).

# A Examples of implementation using use-defined classes

# A.1 Class FullParticle

# A.1.1 Summary

Class FullParticle has all the data of a particle, and a user-defined class given to FDPS with procedure 0 in section 2.3. Users can attach any member variables and functions to this class. However, they need to define several member functions with fixed names. This is because FDPS accesses information of class FullParticle, using these member functions. In what follows, we describe presumptions in this section, necessary member functions, and member functions necessary for some situations.

# A.1.2 Presumptions

In this section, we use class FP as an example of class FullParticle. Users can change the name of FP as they like.

```
class FP;
```

# A.1.3 Necessary member functions

# A.1.3.1 Summary

Necessary member functions are functions FP::getPos and FP::copyFromForce. Using the function FP::getPos, FDPS reads a position from class FullParticle. Using the function FP::copyFromForce, FDPS writes results of interaction calculation to class FullParticle. We present description examples of these functions.

# A.1.3.2 FP::getPos

```
class FP {
public:
    PS::F64vec getPos() const;
};
```

# • Arguments

None

#### • Return vale

Types PS::F32vec or PS::F64vec. A member variable keeping a position of class FP object.

#### • Behavior

Return a member variable keeping a position of class FP object.

# A.1.3.3 FP::copyFromForce

```
class Force {
public:
    PS::F64vec acc;
    PS::F64
               pot;
};
class FP {
public:
    PS::F64vec acceleration;
    PS::F64
               potential;
    void copyFromForce(const Force & force) {
        this->acceleration = force.acc;
        this->potential
                            = force.pot;
    }
};
```

# • Presumptions

Class Force has results of interaction calculation of a particle.

# • Arguments

force: input. Type const Force. The variable force has results of interaction calculation of a particle.

# • Return value

なし。

# • Behavior

Write results of interaction calculation of a particle to class FP object. The member variables acc and pot in class Force correspond to the member variables of acceleration and potential in class FP, respectively.

# • Remarks

The names of "Force" and its member variables are mutable. The names of member variables of class FP are also mutable. The argument name of member function FP::copyFromForce is mutable.

#### A.1.4 Member functions necessary for some situations

#### A.1.4.1 Summary

We describe member functions necessary for some situations. These situations are where all the types of PS::SEARCH\_MODE but type PS::SEARCH\_MODE\_LONG are used, where file I/O APIs of class ParticleSystem are used, where ParticleSystem::adjustPositionIntoRootDomain is used, and where class Particle Mesh is used.

# A.1.4.2 The cases where all the types of PS::SEARCH\_MODE but type PS::SEARCH\_MODE\_LONG are used

# A.1.4.2.1 FP::getRSearch

```
class FP {
public:
    PS::F64 search_radius;
    PS::F64 getRSearch() const {
        return this->search_radius;
    }
};
```

# • Presumptions

A member variable search\_radius is a radius of a particle which is used to search for neighbor particles of a particle. The data type of the search\_radius is types PS::F32 or PS::F64.

# • Arguments

None

#### • Return value

Types PS::F32 or PS::F64. A member variable of class FP object which has a radius of a particle to search for neighbor particles.

#### • Behavior

Return the member variable of class FP object which has a radius of a particle to search for its neighbor particles.

#### • Remarks

The name of member variable search\_radius is mutable.

# A.1.4.3 The case where file I/O APIs of class ParticleSystem is used

Member functions readAscii, writeAscii, readBinary, and writeBinary are necessary when ParticleSystem::readParticleAscii, ParticleSystem::writeParticleAscii, ParticleSystem and ParticleSystem::writeParticleBinary are used, respectively (users can also use different names for these member functions. For more details, please see section 9.1.3.2.3). In what follows, we describe how to describe these member functions.

# A.1.4.3.1 FP::readAscii

#### • Premises

In the input file of particle data, the first, second, third, fourth, and fifth columns contain particle IDs, masses, and three components of positions, respectively. File format is Ascii. 3D Cartesian coordinate system is adopted.

#### • Arguments

fp: type FILE \*. File pointer which points to an input file with particle data.

#### • Return value

None.

#### • Behavior

Read information of id, mass, and pos of class FP from an input file.

#### • Remarks

None.

# A.1.4.3.2 FP::writeAscii

#### • Premises

In the output file of particle data, the first, second, third, fourth, and fifth columns contain particle IDs, masses, and three components of positions, respectively. File format is Ascii. 3D Cartesian coordinate system is adopted.

#### • Arguments

fp: type FILE \*. File pointer which points to an output file with particle data.

#### • Return value

None.

#### • Behavior

Write information of id, mass, and pos of class FP to an output file.

#### • Remarks

None.

# A.1.4.3.3 FP::readBinary

```
class FP {
public:
    PS::S32 id;
    PS::F64 mass;
    PS::F64vec pos;
    void readBinary(FILE *fp) {
        fread(&this->id, sizeof(PS::S32), 1, fp);
        fread(&this->mass, sizeof(PS::F64), 1, fp);
        fread(&this->pos, sizeof(PS::F64vec), 1, fp);
    }
};
```

#### • Premises

In the input file, particle data are assumed to be stored contiguously. In other words, there is no interval between any two particle data. Data of a single particle is stored as follows: particle ID, mass, and three components of position line up in this order. The file position indicator is assumed to point the head of a particle data. File format is binary. 3D Cartesian coordinate system is adopted.

#### • Arguments

fp: type FILE \*. File pointer which points to an input file with particle data.

# • Return value

None.

#### • Behavior

Read information of id, mass, and pos of class FP from an input file.

# • Remarks

None.

# A.1.4.3.4 FP::writeBinary

```
class FP {
public:
    PS::S32 id;
    PS::F64 mass;
    PS::F64vec pos;
    void writeBinary(FILE *fp) {
        fwrite(&this->id, sizeof(PS::S32), 1 , fp);
        fwrite(&this->mass, sizeof(PS::F64), 1, fp);
        fwrite(&this->pos, sizeof(PS::F64vec), 1, fp);
    }
};
```

#### • Premises

In the output file, particle data are assumed to be stored contiguously. In other words, there is no interval between any two particle data. The file position indicator is assumed to point the tail end of the data that is output previously. Now, we want to output a particle data as follows: particle ID, mass, and three components of position line up in this order. File format is binary. 3D Cartesian coordinate system is adopted.

#### • Arguments

fp: type FILE \*. File pointer which points to an output file with particle data.

# • Return value

None.

#### • Behavior

Write information of id, mass, and pos of class FP to an output file.

#### • Remarks

None.

# A.1.4.4 The case where ParticleSystem::adjustPositionIntoRootDomain is used

# A.1.4.4.1 FP::setPos

```
class FP {
public:
    PS::F64vec pos;
    void setPos(const PS::F64vec pos_new) {
        this->pos = pos_new;
    }
};
```

# • Presumptions

The member variable pos is a position of a particle. Its data type is PS::F32vec or PS::F64vec.

# • Arguments

pos\_new: input. Type const PS::F32vec or const PS::F64vec. Position data modified by FDPS.

#### • Return value

None.

#### Behavior

Write position data modified by FDPS to class FP object.

#### • Remarks

The name of member variable pos is mutable. The argument name of pos\_new is mutable. If the data types of pos and pos\_new are different, it may not work well.

#### A.1.4.5 The case where class Particle Mesh is used

When class Particle Mesh is used, member functions FP::getChargeParticleMesh and FP::copyFromForceParticleMesh are necessary. In the following, we describe how to describe them.

# A.1.4.5.1 FP::getChargeParticleMesh

```
class FP {
public:
    PS::F64 mass;
    PS::F64 getChargeParticleMesh() const {
        return this->mass;
    }
};
```

# • Presumptions

The member variable mass has particle mass or charge. Its data type is PS::F32 or PS::F64.

# • Arguments

None.

#### • Return value

Types PS::F32 or PS::F64. Return particle mass or charge.

#### • Behavior

Return a member variable keeping particle mass or charge.

#### • Remarks

The variable name of mass is mutable.

# A.1.4.5.2 FP::copyFromForceParticleMesh

```
class FP {
public:
    PS::F64vec accelerationFromPM;
    void copyFromForceParticleMesh(const PS::F32vec & acc_pm) {
        this->accelerationFromPM = acc_pm;
    }
};
```

# • Presumptions

The member variable accelerationFromPM\_pm has force data exerted on a particle. Its data type is PS::F32vec or PS::F64vec.

# • Arguments

acc\_pm: type const PS::F32vec or const PS::F64vec. Results of class Particle Mesh.

#### • Return value

None.

#### • Behavior

Write a force on a particle from class Particle Mesh to the member variable.

#### • Remarks

The variable name of acc\_pm is mutable. The argument name acc\_pm is mutable.

# A.2 Class EssentialParticleI

# A.2.1 Summary

Class EssentialParticleI has *i*-particle data, and is necessary when users define particle-particle interaction (procedure 0 in section 2.3). The data set of class EssentialParticleI is a subset of the data set of class FullParticle. FDPS accesses the data variables of class EssentialParticleI. In what follows, we describe the presumptions in this section, necessary member functions, and member functions necessary for some situations.

# A.2.2 Presumptions

In this section, we use class EPI and FP as examples of classes EssentialParticleI and Full-Particle, respectively. The names EPI and FP are mutable.

The classes EPI and FP are declared as follows.

```
class FP;
class EPI;
```

# A.2.3 Necessary member functions

# A.2.3.1 Summary

The member functions EPI::getPos and EPI::copyfromFP are necessary. FDPS uses the former to read position data in class EPI, and the latter to write class FP data to class EPI. We present how to describe these functions.

#### A.2.3.2 EPI::getPos

```
class EPI {
  public:
    PS::F64vec pos;
    PS::F64vec getPos() const {
        return this->pos;
    }
};
```

# • Presumptions

The member variable postis particle position. Its data type is PS::F64vec.

#### • Arguments

None.

#### • Return value

Type PS::F64vec. Member variable keeping position data of class EPI.

# • Behavior

Return a memeber variable with position data of class EPI object.

#### • Remarks

The variable name pos is mutable.

# A.2.3.3 EPI::copyFromFP

```
class FP {
public:
    PS::S64
               identity;
    PS::F64
               mass;
    PS::F64vec position;
    PS::F64vec velocity;
    PS::F64vec acceleration;
    PS::F64
               potential;
};
class EPI {
public:
    PS::S64
               id;
    PS::F64vec pos;
    void copyFromFP(const FP & fp) {
        this->id = fp.identity;
        this->pos = fp.position;
    }
};
```

# • Presumptions

The member variable identity and position in class FP correspond to the member variables id and pos in class EPI.

#### • Arguments

fp: input. Type const FP &. Information of class FP object.

#### • Return value

None.

# • Behavior

Write a part of particle data of class FP to class EPI object.

# • Remarks

The member variables of classes FP and EPI are mutable. The argument name of EPI::copyFromFP is mutable. The particle data of class EPI is a subset of the particle data of class FP.

# A.2.4 Member functions necessary for some situations

# A.2.4.1 Summary

In this section, we describe member functions necessary for some situations. The situations are where PS::SEARCH\_MODE\_GATHER and PS::SEARCH\_MODE\_SYMMETRY are adopted for type PS::SEARCH\_MODE.

# A.2.4.2 The case where PS::SEARCH\_MODE\_GATHER and PS::SEARCH\_MODE\_SYMMETRY are adopted for type PS::SEARCH\_MODE

# A.2.4.2.1 EPI::getRSearch

```
class EPI {
public:
    PS::F64 search_radius;
    PS::F64 getRSearch() const {
        return this->search_radius;
    }
};
```

#### • Presumptions

A member variable search\_radius is a radius of a particle which is used to search for neighbor particles of a particle. The data type of the search\_radius is types PS::F32 or PS::F64.

# • Arguments

None.

#### • Return value

Types PS::F32 or PS::F64. A member variable of class EPI object which has a radius of a particle to search for neighbor particles.

# • Behavior

Return the member variable of class EPI object which has a radius of a particle to search for its neighbor particles.

#### • Remarks

The name of member variable search\_radius is mutable.

# A.3 Class EssentialParticleJ

#### A.3.1 Summary

Class EssentialParticle J has j-particle data necessary for interaction calculations, and is necessary for the definition of interactions (procedure 0 in section 2.3). The data of class

EssentialParticleJ is a subset of the data of class FullParticle (section 7.2). FDPS accesses the data of this class. For this purpose, class EssntialParticleJ must have some member functions. In the following, we describe the presumptions of this section, necessary member functions, and member functions necessary for some situations.

# A.3.2 Presumptions

In this section, we use class EPJ and FP as examples of classes EssentialParticleJ and FullParticle, respectively. The class names of EPJ and FP are mutable.

The declarations of EPJ and FP are as follows.

```
class FP; class EPJ;
```

# A.3.3 Necessary member functions

# A.3.3.1 Summary

Necessary member functions are EPJ::getPos and EPJ::copyfromFP. Using the former function, FDPS reads the position data of class FP, and write it to class EPJ. Using the latter function, FDPS reads the various data of class FP, and write them to class EPJ. We describe these functions in this section.

# A.3.3.2 EPJ::getPos

```
class EPJ {
  public:
    PS::F64vec pos;
    PS::F64vec getPos() const {
        return this->pos;
    }
};
```

#### • Presumptions

Member variable pos is the position data of one particle. Its data type is PS::F64vec.

#### • Arguments

None.

#### • Return value

Type PS::F64vec. Member variable keeping position data.

# • Behavior

Return position data of class EPJ object.

# • Remarks

The member variable name of pos is mutable.

# A.3.3.3 EPJ::copyFromFP

```
class FP {
public:
    PS::S64
               identity;
    PS::F64
               mass;
    PS::F64vec position;
    PS::F64vec velocity;
    PS::F64vec acceleration;
    PS::F64
               potential;
};
class EPJ {
public:
    PS::S64
               id;
    PS::F64
               m;
    PS::F64vec pos;
    void copyFromFP(const FP & fp) {
        this->id = fp.identity;
                  = fp.mass;
        this->m
        this->pos = fp.position;
    }
};
```

# • Presumptions

Member variables of identity, mass, and position in class FP correspond to those of id, m, and pos in class EPJ.

# • Arguments

fp: input. Type const FP &. The variable keeping the data of class FP object.

# • Return value

None.

#### • Behavior

Read a part of the data of class FP object, and write them to class EPJ object.

#### • Remarks

The member variable names of classes FP and EPJ are mutable. The argument name of a member function EPJ::copyFromFP is mutable.

# A.3.4 Member functions necessary for some situations

# A.3.4.1 Summary

In this section, we describe member functions necessary for some situations. The situations are where users adopt all but PS::SEARCH\_MODE\_LONG for data type PS::SEARCH\_MODE, and where they adopt all but PS::BOUNDARY\_CONDITION\_OPEN for data type BOUND-ARY\_CONDITION. We do not describe member functions necessary for situations where pre-existing Moment and SuperParticleJ classes. These member functions are explained in sections 7.5 and 7.6.

# A.3.4.2 The cases where all but PS::SEARCH\_MODE\_LONG are used for data type PS::SEARCH\_MODE

# A.3.4.2.1 EPJ::qetRSearch

```
class EPJ {
public:
    PS::F64 search_radius;
    PS::F64 getRSearch() const {
        return this->search_radius;
    }
};
```

#### Presumptions

The member variable search\_radius is a radius of a particle. Neighbor particles of the particle are defined as those inside the particle. The data type of search\_radius may be PS::F32 or PS::F64.

# • Arguments

None.

#### • Return value

Data types PS::F32 or PS::F64. A radius of a particle.

# • Behavior

Return a radius of particle.

#### • Remarks

The member variable name of search\_radius is mutable.

# A.3.4.3 The cases where all but PS::BOUNDARY\_CONDITION\_OPEN are used for data type BOUNDARY\_CONDITION

# A.3.4.3.1 EPJ::setPos

```
class EPJ {
public:
    PS::F64vec pos;
    void setPos(const PS::F64vec pos_new) {
        this->pos = pos_new;
    }
};
```

# • Presumptions

The member variable pos is a position data of a particle. The data type of pos may be PS::F32vec or PS::F64vec. The original data of the variable pos is a member variable of class FP, position. This data type may be PS::F32vec or PS::F64vec.

# • Arguments

pos\_new: Input. Type const PS::F32vec or const PS::F64vec. Position data modified by FDPS.

#### • Return value

None.

#### • Behavior

FDPS writes modified position data to a position data of class EPJ.

#### • Remarks

The member variable name of pos is mutable. The argument name of pos\_new is mutable.

# A.4 Class Moment

#### A.4.1 Summary

Class Moment has moment data of several particles close to each other, and is necessary for the definition of interaction (procedure 0 in section 2.3). Examples of moment data are monopole and dipole of several particles, and the maximum size of these particles. This class plays a role like intermediate variables between classes EssentialParticleJ and SuperParticleJ. Necessary member functions of this class are, for example, a member function which reads some data from class EssnetialParticleJ, and calculate moment of particles.

Some classes are pre-existing. We first describe these classes. Next, we describe how to make class Moment, and what member functions are needed.

# A.4.2 Pre-existing classes

# A.4.2.1 Summary

There are several pre-existing classes. These classes are available when some PS::SEARCH\_MODE types are adopted. In the following, we describe class Moment available when PS::SEARCH\_MODE\_LONG and PS::SEARCH\_MODE\_LONG\_CUTOFF are adopted. When the other PS::SEARCH\_MODE are adopted, users need not to program, considering class Moment.

#### A.4.2.2 PS::SEARCH\_MODE\_LONG

In the Moment classes described below, for MomentMonopole and MomentQuadrupole, their equivalents are defined for PS::SEARCH\_MODE\_LONG\_SCATTER which can be used for the neighbor search. They are MomentMonopoleScatter and MomentQuadrupoleScatter, respectively.

#### A.4.2.2.1 PS::MomentMonopole

This class has moment up to monopole. When this moment is calculated, the center of the coordinate is taken to the center of mass or charge. In the following, we describe an overview of this class.

```
namespace ParticleSimulator {
   class MomentMonopole {
    public:
       F32 mass;
      F32vec pos;
   };
}
```

- Class name PS::MomentMonopole
- Member variables and their information
  mass: the total mass or charge of particles close to each other.
  pos: the center of mass or charge of particles close to each other.
- Use conditions

Class EssentialParticleJ (section 7.4) has member functions EssentialParticleJ::getCharge and EssentialParticleJ::getPos.

In the case of MomentMonopoleScatter, class EssentialParticleJ has the member function getRSearch()

#### A.4.2.2.2 PS::MomentQuadrupole

This class has moment up to quadrupole. When this moment is calculated, the center of the coordinate is taken to the center of mass or charge. In the following, we describe an

overview of this class.

```
namespace ParticleSimulator {
   class MomentQuadrupole {
    public:
        F32 mass;
        F32vec pos;
        F32mat quad;
    };
}
```

- Class name PS::MomentQuadrupole
- Member variables and their information

mass: the total mass of particles close to each other.

pos: the center of mass of particles close to each other.

quad: the quadrupole of mass of particles close to each other.

• Use conditions

Class EssentialParticleJ (section 7.4) has member functions EssentialParticleJ::getCharge and EssentialParticleJ::getPos.

In the case of MomentQuadlepoleScatter, class EssentialParticleJ has the member function getRSearch()

# A.4.2.2.3 PS::MomentMonopoleGeometricCenter

This class has moment up to monopole. When this moment is calculated, the center of the coordinate is taken to the geometric center of particles. In the following, we describe an overview of this class.

- Class name PS::MomentMonopoleGeometricCenter
- Member variables and their information

charge: the total mass or charge of particles close to each other.

pos: the geometric center of particles close to each other.

#### • Use conditions

Class EssentialParticleJ (section 7.4) has member functions EssentialParticleJ::getCharge and EssentialParticleJ::getPos.

# A.4.2.2.4 PS::MomentDipoleGeometricCenter

This class has moment up to dipole. When this moment is calculated, the center of the coordinate is taken to the geometric center of particles. In the following, we describe an overview of this class.

• Class name

PS::MomentDipoleGeometricCenter

• Member variables and their information

charge: the total mass or charge of particles close to each other.

pos: the geometric center of particles close to each other.

dipole: dipole of particle masses or charges.

• Use conditions

Class EssentialParticleJ (section 7.4) has member functions EssentialParticleJ::getCharge and EssentialParticleJ::getPos.

# A.4.2.2.5 PS::MomentQuadrupoleGeometricCenter

This class has moment up to quadrupole. When this moment is calculated, the center of the coordinate is taken to the geometric center of particles. In the following, we describe an overview of this class.

• Class name

PS::MomentQuadrupoleGeometricCenter

• Member variables and their information

charge: the total mass or charge of particles close to each other.

pos: the geometric center of particles close to each other.

dipole: dipole of particle masses or charges.

quadrupole: quadrupole of particle masses or charges.

• Use conditions

Class EssentialParticleJ (section 7.4) has member functions EssentialParticleJ::getCharge and EssentialParticleJ::getPos.

# A.4.2.3 PS::SEARCH\_MODE\_LONG\_CUTOFF

In the Moment classes described below, for MomentMonopoleCutoff, its equivalent is defined for PS::SEARCH\_MODE\_LONG\_CUTOFF\_CATTER which can be used for the neighbor search. It is MomentMonopoleCutoffScatter.

#### A.4.2.3.1 PS::MomentMonopoleCutoff

This class has moment up to monopole. When this moment is calculated, the center of the coordinate is taken to the center of mass or charge. In the following, we describe an overview of this class.

```
namespace ParticleSimulator {
    class MomentMonopoleCutoff {
    public:
        F32 mass;
        F32vec pos;
    };
}
```

• Class name

PS::MomentMonopoleCutoff

• Member variables and their information

mass: the total mass or charge of particles close to each other.

pos: the center of mass or charge of particles close to each other.

• Use conditions

Class EssentialParticleJ (section 7.4) has member functions EssentialParticleJ::getCharge and EssentialParticleJ::getPos.

In the case of MomentMonopoleCutoffScatter, class EssentialParticleJ has the member function getRSearch()

# A.4.3 Necessary member functions

#### A.4.3.1 Summary

In the following, we describe necessary member functions when users define class Moment. As an example, we use Mom for class Moment name. This name is mutable.

# A.4.3.2 Constructor

```
class Mom {
public:
    PS::F32     mass;
    PS::F32vec pos;
    Mom () {
        mass = 0.0;
        pos = 0.0;
    }
};
```

# • Presumptions

Member variables mass and post are mass and position information of class Mom, respectively.

#### • Arguments

None.

#### • Return value

None.

#### • Behavior

Initialize class Mom object.

# • Remarks

The member variable names are mutable. Some member variables can be added.

# • Presumptions

Member variables mass and post are mass and position information of class Mom, respectively.

# • Arguments

```
m: input. Type const PS::F32. Mass.p: Input. Type const PS::F32vec &. Position.
```

# • Return value

None.

#### • Behavior

Initialize class Mom object.

# • Remarks

The member variable names are mutable. Some member variables can be added.

#### A.4.3.3 Mom::init

```
class Mom {
  public:
    void init();
};
```

# • Presumptions

None.

# • Arguments

None.

# • Return value

None.

#### • Behavior

Initialize class Mom object.

#### • Remarks

None.

# A.4.3.4 Mom::getPos

```
class Mom {
public:
    PS::F32vec pos;
    PS::F32vec getPos() const {
        return pos;
    }
};
```

# • Presumptions

A member variable pos is a representative position of particles close to each other. This data type is PS::F32vec or PS::F64vec.

# • Arguments

None.

#### • Return value

PS::F32vec or PS::F64vec. The member variable pos.

#### • Behavior

Return the member variable pos.

#### • Remarks

The member variable name is mutable.

# A.4.3.5 Mom::getCharge

```
class Mom {
public:
    PS::F32 mass;
    PS::F32 getCharge() const {
        return mass;
    }
};
```

# • Presumptions

A member variable mass is the total mass or charge of particles close to each other. This data type is PS::F32 or PS::F64.

# • Arguments

None.

#### • Return value

Type PS::F32 or PS::F64. The member variable mass.

#### • Behavior

Return the member variable mass.

#### • Remarks

The member variable name is mutable.

#### A.4.3.6 Mom::accumulateAtLeaf

```
class Mom {
public:
    PS::F32    mass;
    PS::F32vec pos;
    template <class Tepj>
    void accumulateAtLeaf(const Tepj & epj) {
        mass += epj.getCharge();
        pos += epj.getPos();
    }
};
```

# • Presumptions

A member variable mass is the total mass or charge of particles close to each other. This data type is PS::F32 or PS::F64. A member variable pos is a representative position of particles close to each other. This data type is PS::F32vec or PS::F64vec. Class EssentialParticleJ is inserted into the template argument Tepj, and has member functions getCharge and getPos.

#### • Arguments

epj: input. Type const Tepj &. An object of class Tpj.

# • Return value

None.

#### Behavior

Calculate moment from an object of class EssentialParticleJ.

#### • Remarks

The member variable names mass and pos are mutable. The argument name epj is mutable.

#### A.4.3.7 Mom::accumulate

```
class Mom {
public:
    PS::F32     mass;
    PS::F32vec pos;
    void accumulate(const Mom & mom) {
        mass += mom.mass;
        pos += mom.mass * mom.pos;
    }
};
```

# • Presumptions

A member variable mass is the total mass or charge of particles close to each other. This data type is PS::F32 or PS::F64. A member variable pos is the center of mass of particles close to each other. This data type is PS::F32vec or PS::F64vec. Class EssentialParticleJ is inserted into the template argument Tepj, and has member functions getCharge and getPos.

# • Arguments

mom: input. Type const Mom &. An object of class Mom.

# • Return value

None.

#### • Behavior

Calculate information of class Mom from an object of class Mom.

#### • Remarks

The member variable names mass and pos are mutable. The argument name epj is mutable.

#### A.4.3.8 Mom::set

```
class Mom {
public:
    PS::F32     mass;
    PS::F32vec pos;
    void set() {
        pos = pos / mass;
    }
};
```

# • Presumptions

A member variable mass is the total mass or charge of particles close to each other. This data type is PS::F32 or PS::F64. A member variable pos is the center of mass of particles close to each other. This data type is PS::F32vec or PS::F64vec. Class EssentialParticleJ is inserted into the template argument Tepj, and has member functions getCharge and getPos.

# • Arguments

None.

#### • Return value

None.

#### • Behavior

Normalize position data of moment, since they are not normalized in the member functions in Mom::accumulateAtLeaf and Mom::accumulate.

# • Remarks

The member variable names mass and pos are mutable. The argument name epj is mutable.

#### A.4.3.9 Mom::accumulateAtLeaf2

```
class Mom {
public:
    PS::F32
               mass;
    PS::F32vec pos;
    PS::F32mat quad;
    template <class Tepj>
    void accumulateAtLeaf2(const Tepj & epj) {
        PS::F64 ctmp
                        = epj.getCharge();
        PS::F64vec ptmp = epj.getPos() - pos;
        PS::F64 cx = ctmp * ptmp.x;
        PS::F64 cy = ctmp * ptmp.y;
        PS::F64 cz = ctmp * ptmp.z;
        quad.xx += cx * ptmp.x;
        quad.yy += cy * ptmp.y;
        quad.zz += cz * ptmp.z;
        quad.xy += cx * ptmp.y;
        quad.xz += cx * ptmp.z;
        quad.yz += cy * ptmp.z;
    }
};
```

#### • Presumptions

A member variable mass is the total mass or charge of particles close to each other. This data type is PS::F32 or PS::F64. A member variable pos is a representative position of particles close to each other. This data type is PS::F32vec or PS::F64vec. This has been already obtained in the member function Mom::accumulateAtLeaf. A member variable quad is quadrupole of particles close to each other. This data type is PS::F32mat or PS::F64mat. Class EssentialParticleJ is inserted into the template argument Tepj, and has member functions getCharge and getPos.

#### • Arguments

epj: input. Type const Tepj &. An object of class Tepj.

#### • Return value

None.

#### Behavior

Calculate moment from an object of class EssentialParticleJ.

#### • Remarks

The member variable names mass, pos and quad are mutable. The argument name epj is mutable.

#### A.4.3.10 Mom::accumulate2

```
class Mom {
public:
    PS::F32
               mass;
    PS::F32vec pos;
    PS::F32mat quad;
    void accumulate(const Mom & mom) {
        PS::F64 mtmp
                        = mom.mass;
        PS::F64vec ptmp = mom.pos - pos;
        PS::F64 cx = mtmp * ptmp.x;
        PS::F64 cy = mtmp * ptmp.y;
        PS::F64 cz = mtmp * ptmp.z;
        quad.xx += cx * ptmp.x + mom.quad.xx;
        quad.yy += cy * ptmp.y + mom.quad.yy;
        quad.zz += cz * ptmp.z + mom.quad.zz;
        quad.xy += cx * ptmp.y + mom.quad.xy;
        quad.xz += cx * ptmp.z + mom.quad.xz;
        quad.yz += cy * ptmp.z + mom.quad.yz;
    }
};
```

# • Presumptions

A member variable mass is the total mass or charge of particles close to each other. This data type is PS::F32 or PS::F64. A member variable pos is a representative position of particles close to each other. This data type is PS::F32vec or PS::F64vec. This has been already obtained in the member function Mom::accumulate. A member variable quad is quadrupole of particles close to each other. This data type is PS::F32mat or PS::F64mat.

# • Arguments

mom: input. Type const Mom &. An object of class Mom.

#### • Return value

None.

#### • Behavior

Calculate information of class Mom from an object of class Mom

#### • Remarks

The member variable names mass, pos and quad are mutable.

# A.5 Class SuperParticleJ

# A.5.1 Summary

Class SuperPrticleJ has information of superparticle which is a representative of particles close to each other. This is required for the definition of interaction (procedure 0 in section 2.3). This class is necessary only when PS::SEARCH\_MODE\_LONG and PS::SEARCH\_MODE\_LONG\_CUTOFF are adopted for type PS::SEARCH\_MODE. This class has member functions communicating position data with FDPS. Information of classes SuperParticleJ and Moment pairs up. Therefore, this class has member functions which transform from data of class Moment to that of class SuperParticleJ, and vice versa.

Some classes are pre-existing. We first describe these classes. Next, we describe how to make class SuperParticleJ, and what member functions are needed.

### A.5.2 Pre-existing classes

There are several pre-existing classes. These classes are available when some PS::SEARCH\_MODE types are adopted. In the following, we describe class SuperParticleJ available when PS::SEARCH\_MODE\_LONG and PS::SEARCH\_MODE\_LONG\_CUTOFF are adopted. When the other PS::SEARCH\_MODE are adopted, users need not to program, considering class SuperParticleJ.

#### A.5.2.1 PS::SEARCH\_MODE\_LONG

# A.5.2.1.1 PS::SPJMonopole

This class has moment up to monopole, and is paired with class PS::MomentMonopole. We overview this class as follows.

```
namespace ParticleSimulator {
   class SPJMonopole {
    public:
        F64 mass;
        F64vec pos;
    };
}
```

- Class name PS::SPJMonopole
- Member variables and their information
  mass: the total mass or charge of particles close to each other.
   pos: the center of mass or charge of particles close to each other.
- Use conditions

The same as class PS::MomentMonopole.

# A.5.2.1.2 PS::SPJQuadrupole

This class has monopole and quadrupole moments, and is paired with class PS::MomentQuadrupole. In the following, we describe an overview of this class.

```
namespace ParticleSimulator {
   class SPJQuadrupole {
    public:
        F32 mass;
        F32vec pos;
        F32mat quad;
    };
}
```

- Class name PS::SPJQuadrupole
- Member variables and their information

mass: the total mass of particles close to each other.

pos: the center of mass of particles close to each other.

quad: the quadrupole of mass of particles close to each other.

• Use conditions

The same as class PS::MomentQuadrupole.

# A.5.2.1.3 PS::SPJMonopoleGeometricCenter

This class has moment up to monopole, and is paired with class PS::MomentMonpoleGeometricCenter. In the following, we describe an overview of this class.

• Class name

PS::SPJMonopoleGeometricCenter

• Member variables and their information

charge: the total mass or charge of particles close to each other.

pos: the geometric center of particles close to each other.

• Use conditions

The same as class PS::MomentMonopoleGeometricCenter.

# A.5.2.1.4 PS::SPJDipoleGeometricCenter

This class has moment up to dipole, and is paired with class PS::MomentDipoleGeometricCenter. In the following, we describe an overview of this class.

• Class name

PS::SPJDipoleGeometricCenter

• Member variables and their information

charge: the total mass or charge of particles close to each other.

pos: the geometric center of particles close to each other.

dipole: dipole of particle masses or charges.

• Use conditions

The same as class PS::MomentDipoleGeometricCenter.

# A.5.2.1.5 PS::SPJQuadrupoleGeometricCenter

This class has moment up to quadrupole, and is paired with class PS::MomentQuadrupoleGeometricCenter. In the following, we describe an overview of this class.

• Class name

PS::SPJQuadrupoleGeometricCenter

• Member variables and their information

charge: the total mass or charge of particles close to each other.

pos: the geometric center of particles close to each other.

dipole: dipole of particle masses or charges.

quadrupole: quadrupole of particle masses or charges.

• Use conditions

The same as class PS::MomentQuadrupoleGeometricCenter.

#### A.5.2.2 PS::SEARCH\_MODE\_LONG\_CUTOFF

#### A.5.2.2.1 PS::SPJMonopoleCutoff

This class has moment up to monopole, and is paired with class PS::MomentMonopoleCutoff. In the following, we describe an overview of this class.

```
namespace ParticleSimulator {
   class SPJMonopoleCutoff {
    public:
       F32 mass;
      F32vec pos;
   };
}
```

• Class name

PS::SPJMonopoleCutoff

• Member variables and their information

mass: the total mass or charge of particles close to each other.

pos: the center of mass or charge of particles close to each other.

• Use conditions

The same as class PS::MomentMonopoleCutoff.

# A.5.3 Necessary member functions

# A.5.3.1 Summary

We describe necessary member functions when users make class SuperParticleJ. We use class SPJ for an example of class SuperParticleJ. This name is mutable.

# A.5.3.2 SPJ::getPos

```
class SPJ {
public:
    PS::F64vec pos;
    PS::F64vec getPos() const {
        return this->pos;
    }
};
```

# • Presumptions

Member variables pos is position information of this class. This data type is PS::F32vec or PS::F64vec.

# • Arguments

None.

#### • Return value

Type PS::F32vec or PS::F64vec. Member variable keeping position data of class SPJ.

#### • Behavior

Return position data of an object of class SPJ.

#### • Remarks

The member variable name is mutable.

#### A.5.3.3 SPJ::setPos

```
class SPJ {
  public:
    PS::F64vec pos;
    void setPos(const PS::F64vec pos_new) {
        this->pos = pos_new;
    }
};
```

# • Presumptions

A member variable pos is position data of a particle. This data type is PS::F32vec or PS::F64vec.

# • Arguments

pos\_new: input. Type const PS::F32vec or const PS::F64vec. Position modified by FDPS.

## • Return value

None.

### • Behavior

Write position data modified by FDPS to position data of an object of class SPJ.

### • Remarks

The member variable name is mutable. The argument name is mutable.

# A.5.3.4 SPJ::copyFromMoment

```
class Mom {
public:
    PS::F32    mass;
    PS::F32vec pos;
}
class SPJ {
public:
    PS::F32    mass;
    PS::F32vec pos;
    void copyFromMoment(const Mom & mom) {
        mass = mom.mass;
        pos = mom.pos;
    }
};
```

## • Presumptions

None.

# • Arguments

mom: input. Type const Mom &. User-defined class Moment.

## • Return value

None.

## • Behavior

Copy an object of class Mom to that of class SPJ.

### • Remarks

Class name Mom is mutable. Member variables of classes Mom and SPJ are mutable. The argument name of the function SPJ::copyFromMoment is mutable.

# A.5.3.5 SPJ::convertToMoment

```
class Mom {
public:
    PS::F32
               mass;
    PS::F32vec pos;
    Mom(const PS::F32 m,
        const PS::F32vec & p) {
        mass = m;
        pos = p;
class SPJ {
public:
    PS::F32
               mass;
    PS::F32vec pos;
    Mom convertToMoment() const {
        return Mom(mass, pos);
    }
};
```

# • Presumptions

None.

## • Arguments

None.

## • Return value

Type Mom. Constructor of class Mom.

### • Behavior

Return the constructor of class Mom.

## • Remarks

Class name Mom is mutable. Member variable names of classes Mom and SPJ are mutable. The argument name of the function SPJ::copyFromMoment is mutable. Predefinition of the constructor of class Mom is necessary.

## A.5.3.6 SPJ::clear

```
class SPJ {
public:
    PS::F32    mass;
    PS::F32vec pos;
    void clear() {
        mass = 0.0;
        pos = 0.0;
    }
};
```

## • Presumptions

None.

# • Arguments

None.

## • Return value

None.

### • Behavior

Clear information of an object of class SPJ.

### • Remarks

Member variable names are mutable.

# A.6 Class Force

# A.6.1 Summary

This class has results of interaction calculations, and is necessary for the definition of interaction (procedure 0 in section 2.3). We describe presumptions in this section, and necessary member functions.

# A.6.2 Presumptions

We use class Result as an example of class Force. This name is mutable.

# A.6.3 Necessary member functions

A necessary member function is Result::clear. This function initializes results of interaction calculations.

## A.6.3.1 Result::clear

```
class Result {
public:
    PS::F32vec acc;
    PS::F32    pot;
    void clear() {
        acc = 0.0;
        pot = 0.0;
    }
};
```

# • Presumptions

Member variables are acc and pot.

# • Arguments

None.

## • Return value

None.

### • Behavior

Initialize member variables.

### • Remarks

The member variable names are mutable.

# A.7 Class Header

# A.7.1 Summary

Class Header defines a format of a header of I/O files. This class is used in file I/O APIs of class ParticleSystem. These APIs are ParticleSystem::readParticleAscii and ParticleSystem::writeParticleAscii. We present presumptions in this section and member functions necessary when users use these APIs.

### A.7.2 Presumptions

We use class Hdr as an example of class Header. This name is mutable.

# A.7.3 Member functions necessary for some situations

### A.7.3.1 Hdr::readAscii

```
class Hdr {
public:
    PS::S32 nparticle;
    PS::F64 time;
    PS::S32 readAscii(FILE *fp) {
        fscanf(fp, "%d%lf", &this->nparticle, &this->time);
        return this->nparticle;
    }
};
```

### • Presumptions

This header has the number of particles "nparticle" and time "time".

### • Arguments

fp: input. Type FILE \*. File pointer to an input file.

### • Return value

Type PS::S32. Return the number of particles, and return -1 unless there is the number of particles in the header.

### • Behavior

Read the header information from the input file of particle data.

## • Remarks

The member variable names are mutable. It is undefined when the return value is neither the number of particles and -1.

### A.7.3.2 Hdr::writeAscii

```
class Hdr {
public:
    PS::S32 nparticle;
    PS::F64 time;
    void writeAscii(FILE *fp) {
        fprintf(fp, "%d %lf", this->nparticle, this->time);
    }
};
```

## • Presumptions

This header has the number of particles "nparticle" and time "time".

# • Arguments

fp: input. Type FILE \*. File pointer to an output file.

## • Return value

None.

## • Behavior

Write the header information to the output file.

### • Remarks

The member variable names are mutable.

# A.8 Function object calcForceEpEp

## A.8.1 Summary

Function object calcForceEpEp presents an interaction, and is necessary for the definition of the interaction (procedure 0 in section 2.3). In what follows, we describe how to describe this.

# A.8.2 Presumptions

We show the function object for calculating gravitational N-body simulation. The object name is gravityEpEp, and is mutable. We use classes EPI and EPJ for classes EssentialParitleeI and EssentialParitleeJ, respectively.

# A.8.3 gravityEpEp::operator ()

Listing 47: calcForceEpEp

```
1 class Result {
2 public:
3
       PS::F32vec acc;
4 };
5 class EPI \{
  public:
7
       PS:: S32
                    id;
       PS::F32vec pos;
8
9 };
10 class EPJ {
11 public:
12
       PS:: S32
                    id;
13
       PS::F32
                    mass;
       PS::F32vec pos;
14
15 };
16 struct gravityEpEp {
17
       static PS::F32 eps2;
```

```
18
       void operator () (const EPI *epi,
19
                           const PS::S32 ni,
20
                           const EPJ *epj,
21
                           const PS::S32 nj,
22
                           Result *result) {
23
24
           for (PS::S32 i = 0; i < ni; i++) {
25
                PS::S32
                            ii = epi[i].id;
26
                PS::F32vec xi = epi[i].pos;
27
                PS::F32vec ai = 0.0;
28
                for (PS::S32 j = 0; j < nj; j++) {
29
                    PS::S32
                                jj = epj[j].id;
30
                    PS::F32
                                mj = epj[j].mass;
31
                    PS::F32vec xj = epj[j].pos;
32
33
                    PS::F32vec dx
                                      = xi - xj;
34
                    PS::F32
                                r2
                                      = dx * dx + eps2;
35
                    PS::F32
                                rinv = (ii != jj) ? 1. / sqrt(r2)
36
                                                     : 0.0;
37
38
                    ai += mj * rinv * rinv * rinv * dx;
39
                }
40
                result.acc = ai;
41
           }
       }
42
43 };
44 PS::F32 gravityEpEp::eps2 = 9.765625e-4;
```

### • Presumptions

Member functions necessary for classes Result, EPI, and EPJ are omitted. A member variable acc in class Result is gravitational acceleration from j-particle to i-particle. Member variables id and pos in classes EPI and EPJ are particle ID and position, respectively. A member variable mass in class EPJ is mass of j-particle. A member variable eps2 in function object gravityEpEp is square of gravitational softening. Users need not to consider multi-threading, since multi-threading is defined outside this function object.

# • Arguments

```
epi: input. Type const EPI * or EPI *. Array of i-particle data. ni: input. Type const PS::S32 or PS::S32. The number of i-particles. epj: input. Type const EPJ * or EPJ *. Array of j-particle data. nj: input. Type const PS::S32 or PS::S32. The number of j-particles. result: output. Type Result *. Array of results of interactions of i-particles.
```

### • Return value

None.

#### Behavior

Calculate forces exerted by j-particles on i-particles.

### • Remarks

All the argument names are mutable. The contents of the function object are mutable.

# A.9 Function object calcForceSpEp

# A.9.1 Summary

Function object calcForceSpEp presents forces exerted by superparticles on particles, and is necessary for the definition of the interaction (procedure 0 in section 2.3). In what follows, we describe how to describe this.

# A.9.2 Presumptions

We show the function object for calculating gravitational N-body simulation. The object name is gravitySpEp, and is mutable. We use classes EPI and SPJ for classes EssentialParitleeI and SuperParitleeJ, respectively.

# A.9.3 gravitySpEp::operator ()

Listing 48: calcForceSpEp

```
1 class Result {
  public:
3
       PS::F32vec accfromspj;
4 };
  class EPI {
  public:
7
       PS:: S32
                   id:
8
       PS::F32vec pos;
9 };
10 class SPJ {
11 public:
12
       PS::F32
                   mass;
13
       PS::F32vec pos;
14 };
15
  struct gravitySpEp {
       static PS::F32 eps2;
16
17
       void operator () (const EPI *epi,
18
                           const PS::S32 ni,
19
                           const SPJ *spj,
20
                           const PS::S32 nj,
```

```
21
                            Result *result) {
22
23
            for(PS::S32 i = 0; i < ni; i++) {
24
                PS::F32vec xi = epi[i].pos;
25
                PS::F32vec ai = 0.0;
26
                for (PS::S32 j = 0; j < nj; j++) {
27
                                 mj = spj[j].mass;
                     PS::F32
28
                     PS::F32vec xj = spj[j].pos;
29
                                       = xi - xj;
30
                     PS::F32vec dx
31
                     PS::F32
                                 r2
                                       = dx * dx + eps2;
32
                     PS::F32
                                 rinv = 1. / sqrt(r2);
33
34
                     ai += mj * rinv * rinv * rinv * dx;
35
                }
36
                result.accfromspj = ai;
37
            }
38
       }
39 };
40 \text{ PS}::F32 \text{ gravitySpEp}::eps2 = 9.765625e-4;
```

# • Presumptions

Member functions necessary for classes Result, EPI, and SPJ are omitted. A member variable accfromspj in class Result is gravitational acceleration from superparticle to *i*-particle. Member variables pos in classes EPI and SPJ are particle position. A member variable mass in class SPJ is mass of superparticle. A member variable eps2 in function object gravitySpEp is square of gravitational softening. Users need not to consider multi-threading, since multi-threading is defined outside this function object.

### • Arguments

```
epi: input. Type const EPI * or EPI *. Array of i-particle data. ni: input. Type const PS::S32 or PS::S32. The number of i-particles. spj: input. Type const SPJ * or SPJ *. Array of superparticle data. nj: input. Type const PS::S32 or PS::S32. The number of superparticles. result: output. Type Result *. Array of results of interactions of i-particles.
```

### • Return value

None.

#### • Behavior

Calculate forces exerted by superparticles on *i*-particles.

### • Remarks

All the argument names are mutable. The contents of the function object are mutable.

# A.10 Functor calcForceDispatch

# A.10.1 Summary

Functor calcForceDispatch is used in the case when some accelerator hardware is used for the interaction calculation. It sends the particles to the accelerator and startup the interaction calculation kernel on the accelerator. In the following, we present an example implementation of this function.

### A.10.2 Premises

Here we present one example of using Cuda to implement gravitational interaction between particles. The name of functor calcForceDispatch is CalcForceDispatch. The names for classes EssentialParitlceI, SuperParitlceJ, and Force are EPI, SPJ, and Result, respectively. These name can be changed to any names legal within C++ grammer.

# A.10.3 The example

Listing 49: calcForceDispatch

```
1
2
  class EpiGPU{
3
  public:
       float2 pos[3];
4
5
       int id_walk;
6
  };
7
8
  class EpjGPU{
9
  public:
10
       float mass;
11
       float2 pos[3];
12 };
13
14 class ForceGPU{
15 public:
16
       float2 acc[3];
17
       float2 pot;
18 };
19
20
  __global__ void ForceKernel(const EpiGPU * epi,
21
                                  const EpjGPU * epj,
22
                                  const int
                                                * nj_disp,
23
                                  ForceGPU
                                                * force,
24
                                  const float eps2){
25
       int id_i = blockDim.x * blockIdx.x + threadIdx.x;
26
       const EpiGPU & ip = epi[id_i];
27
       float2 poti;
28
       float2 acci[3];
```

```
29
       poti = acci[0] = acci[1] = acci[2] = make_float2(0.0, 0.0);
30
       const int j_head = nj_disp[ip.id_walk];
       const int j_tail = nj_disp[ip.id_walk+1];
31
32
       const int nj = j_tail - j_head;
33
       for(int j=j_head; j<j_tail; j++){</pre>
34
           EpjGPU jp = epj[j];
           const float dx = (jp.pos[0].x - ip.pos[0].x) + (jp.pos
35
                 [0].y - ip.pos[0].y);
           const float dy = (jp.pos[1].x - ip.pos[1].x) + (jp.pos
36
                 [1].y - ip.pos[1].y);
           const float dz = (jp.pos[2].x - ip.pos[2].x) + (jp.pos
37
                 [2].y - ip.pos[2].y);
38
           const float r2 = ((eps2 + dx*dx) + dy*dy) + dz*dz;
39
           const float r_inv = rsqrtf(r2);
40
           const float pij = jp.mass * r_inv * (r2 > eps2);
           const float r2_inv = r_inv * r_inv;
41
42
           const float pij_r3_inv = pij * r2_inv;
43
           const float ax = pij_r3_inv * dx;
44
           const float ay = pij_r3_inv * dy;
45
           const float az = pij_r3_inv * dz;
           poti = float2_accum(poti, pij);
46
47
           acci[0] = float2_accum(acci[0], ax);
48
           acci[1] = float2_accum(acci[1], ay);
49
           acci[2] = float2_accum(acci[2], az);
50
       }
51
       poti = float2_regularize(poti);
52
       acci[0] = float2_regularize(acci[0]);
53
       acci[1] = float2_regularize(acci[1]);
54
       acci[2] = float2_regularize(acci[2]);
55
       force[id_i].pot = poti;
56
       force[id_i].acc[0] = acci[0];
       force[id_i].acc[1] = acci[1];
57
       force[id_i].acc[2] = acci[2];
58
59 }
60
61 static ForceGPU * force_d;
62 static ForceGPU * force_h;
63 static EpiGPU * epi_d;
64 static EpiGPU * epi_h;
65 static EpjGPU * epj_d;
66 static EpjGPU * epj_h;
67 static int * ni_disp_h;
68 static int * nj_disp_d;
69 static int * nj_disp_h;
70
71 int DispatchKernelWithSP(const PS::S32 tag,
```

```
72
                              const int
                                           n_walk,
73
                              const EPIGrav ** epi,
74
                              const int
                                        * n_epi,
75
                              const EPJGrav ** epj,
76
                              const int * n_epj,
77
                              const PS::SPJMonopole ** spj,
78
                              const int
                                         *
                                             n_spj){
79
        static bool first = true;
        assert(n_walk <= N_WALK_LIMIT);</pre>
80
81
       if(first){
82
            CUDA_SAFE_CALL( cudaMalloc(
                                           (void**)&nj_disp_d,
                                                                    (
                  N_WALK_LIMIT+1)*sizeof(int) );
83
            CUDA_SAFE_CALL( cudaMallocHost( (void**)&ni_disp_h,
                                                                    (
                  N_WALK_LIMIT+1)*sizeof(int) );
84
            CUDA_SAFE_CALL( cudaMallocHost( (void**)&nj_disp_h,
                  N_WALK_LIMIT+1)*sizeof(int) );
85
            CUDA_SAFE_CALL( cudaMalloc( (void**)&epi_d,
                  NI_LIMIT*sizeof(EpiGPU) ) );
            CUDA_SAFE_CALL( cudaMalloc( (void**)&epj_d,
86
                  NJ_LIMIT*sizeof(EpjGPU) );
            CUDA_SAFE_CALL( cudaMalloc( (void**)&force_d,
87
                  NI_LIMIT*sizeof(ForceGPU) ) );
88
            CUDA_SAFE_CALL( cudaMallocHost( (void**)&epi_h,
                  NI_LIMIT*sizeof(EpiGPU) ) );
89
            CUDA_SAFE_CALL( cudaMallocHost( (void**)&epj_h,
                  NJ_LIMIT*sizeof(EpjGPU) );
90
            CUDA_SAFE_CALL( cudaMallocHost( (void**)&force_h,
                  NI_LIMIT*sizeof(ForceGPU) ) );
91
            first = false;
92
       }
93
       const float eps2 = EPIGrav::eps * EPIGrav::eps;
94
       ni_disp_h[0] = nj_disp_h[0] = 0;
95
       for(int i=0; i<n_walk; i++){</pre>
96
            ni_disp_h[i+1] = ni_disp_h[i] + n_epi[i];
97
            nj_disp_h[i+1] = nj_disp_h[i] + n_epj[i] + n_spj[i];
98
       }
99
        int ni_tot = ni_disp_h[n_walk];
100
        const int ni_tot_reg = ni_disp_h[n_walk] + ( (ni_tot%
             N_THREAD_GPU != 0) ? (N_THREAD_GPU - (ni_tot%
             N_THREAD_GPU)) : 0);
101
        assert(ni_tot_reg <= NI_LIMIT);</pre>
        assert(nj_disp_h[n_walk] <= NJ_LIMIT);</pre>
102
103
       ni_tot = 0;
       int nj_tot = 0;
104
105
       for(int iw=0; iw<n_walk; iw++){</pre>
106
            for(int ip=0; ip<n_epi[iw]; ip++){</pre>
```

```
107
                epi_h[ni_tot].pos[0] = float2_split(epi[iw][ip].
                      pos.x);
108
                epi_h[ni_tot].pos[1] = float2_split(epi[iw][ip].
                      pos.y);
109
                epi_h[ni_tot].pos[2] = float2_split(epi[iw][ip].
                      pos.z);
                epi_h[ni_tot].id_walk = iw;
110
                force_h[ni_tot].acc[0] = force_h[ni_tot].acc[1]
111
                    = force_h[ni_tot].acc[2] = force_h[ni_tot].pot
112
                          = make_float2(0.0, 0.0);
113
                ni_tot++;
114
115
            for (int jp=0; jp<n_epj[iw]; jp++) {
116
                epj_h[nj_tot].mass
                                    = epj[iw][jp].mass;
117
                epj_h[nj_tot].pos[0] = float2_split(epj[iw][jp].
                      pos.x);
118
                epj_h[nj_tot].pos[1] = float2_split(epj[iw][jp].
                     pos.y);
119
                epj_h[nj_tot].pos[2] = float2_split(epj[iw][jp].
                     pos.z);
120
                nj_tot++;
121
            }
122
            for (int jp=0; jp<n_spj[iw]; jp++) {
                epj_h[nj_tot].mass
123
                                       = spj[iw][jp].getCharge();
124
                epj_h[nj_tot].pos[0] = float2_split(spj[iw][jp].
                      getPos().x);
125
                epj_h[nj_tot].pos[1]
                                     = float2_split(spj[iw][jp].
                      getPos().y);
                epj_h[nj_tot].pos[2]
126
                                     = float2_split(spj[iw][jp].
                      getPos().z);
127
                nj_tot++;
            }
128
129
130
       for(int ip=ni_tot; ip<ni_tot_reg; ip++){</pre>
            epi_h[ni_tot].pos[0] = epi_h[ni_tot].pos[1] = epi_h[
131
                 ni_tot].pos[2] = make_float2(0.0, 0.0);
            epi_h[ni_tot].id_walk = 0;
132
133
            force_h[ni_tot].acc[0] = force_h[ni_tot].acc[1]
134
                = force_h[ni_tot].acc[2] = force_h[ni_tot].pot =
                     make_float2(0.0, 0.0);
135
       }
       CUDA_SAFE_CALL( cudaMemcpy(epi_d, epi_h, ni_tot_reg*sizeof(
136
             EpiGPU), cudaMemcpyHostToDevice) );
137
       CUDA_SAFE_CALL( cudaMemcpy(epj_d, epj_h, nj_tot*sizeof(
             EpjGPU), cudaMemcpyHostToDevice) );
```

```
138
       CUDA_SAFE_CALL( cudaMemcpy(nj_disp_d, nj_disp_h, (n_walk+1)
             *sizeof(int), cudaMemcpyHostToDevice));
        const int n_grid = ni_tot_reg/N_THREAD_GPU + ((ni_tot_reg%)
139
             N_THREAD_GPU == 0) ? 0 : 1);
140
       dim3 size_grid(n_grid, 1, 1);
141
       dim3 size_thread(N_THREAD_GPU, 1, 1);
142
       ForceKernel <<<size_grid, size_thread>>> (epi_d, epj_d,
             nj_disp_d, force_d, float(eps2));
143
144
       return 0;
145 }
```

### • Arguments

tag: input. Type const PS::S32. Corresponding call to CalcForceRetrieve() should use the same value for tag

nwalk: input.Type const PS::S32. The number of interaction lists.

epi: input. Type const EPI\*\* or EPI\*\*. Array of the array of i-particles.

ni: input. Type const PS::S32\* or PS::S32\*. Array of the numbers of i-particles.

spj: input. Type const EPJ\*\* or EPJ\*\*. Array of the array of j-particles.

nj: input. Type const PS::S32\* or PS::S32\*. Array of the number of j-paricles.

• return value Returns 0 upon normal completion. Otherwize non-zero values are returned.

# • Function

Send epi and epj to the accelerator and let the accelerator do the interaction calculation.

#### • Remarks

All the argument names are mutable. The contents of the function object are mutable.

## A.11 Functor calcForceRetrieve

# A.11.1 Summary

Functor calcForceRetrieve retrieves the results calculated on the accelerator. In the following, we present an example implementation of this function.

#### A.11.2 Premises

Here we present one example of using Cuda to implement gravitational interaction between particles. The name of functor calcForceRetrieve is RetieveKernel. The name for the class Force is ForceGrav. These name can be changed to any names legal within C++ grammer.

# Listing 50: calcForceRetrieve

```
int RetrieveKernel(const PS::S32 tag,
1
2
                       const PS::S32
                                          n_walk,
3
                       const PS::S32 *
                                          ni.
4
                       ForceGrav
                                       ** force){
5
       int ni_tot = 0;
6
       for(int i=0; i<n_walk; i++){</pre>
7
           ni_tot += ni[i];
8
9
       CUDA_SAFE_CALL( cudaMemcpy(force_h, force_d,
             sizeof(ForceGPU), cudaMemcpyDeviceToHost) );
10
       int n_cnt = 0;
11
       for(int iw=0; iw<n_walk; iw++){</pre>
           for(int ip=0; ip<ni[iw]; ip++){</pre>
12
13
                force[iw][ip].acc.x = (double)force_h[n_cnt].acc
                      [0].x + (double)force_h[n_cnt].acc[0].y;
                force[iw][ip].acc.y = (double)force_h[n_cnt].acc
14
                      [1].x + (double)force_h[n_cnt].acc[1].y;
15
                force[iw][ip].acc.z = (double)force_h[n_cnt].acc
                      [2].x + (double)force_h[n_cnt].acc[2].y;
                force[iw][ip].pot = (double)force_h[n_cnt].pot.x
16
                         + (double)force_h[n_cnt].pot.y;
17
                force [iw] [ip] . pot *= -1.0;
18
                n_cnt++;
19
           }
20
       }
21
       return 0;
22 }
```

### • Arguments

tag: input. Type const PS::S32. Corresponding call to CalcForceDispatch() should use the same value for tag

nwalk: input.Type const PS::S32. The number of interaction lists.

ni: input. Type const PS::S32\* or PS::S32\*. Array of the numbers of i-particles.

force: output. Type Result\*\*.

• **return value** Returns 0 upon normal completion. Otherwize non-zero values are returned.

# • Function

Store the results calculated on the accielerator to the array force.

#### • Remarks

All the argument names are mutable. The contents of the function object are mutable.