



Materials and geometry

Giada Petringa (LNS-INFN)

*The 2nd Geant4 School in China
Shandong University
Qingdao*

User classes (starting...)

At initialization

G4VUserDetectorConstruction

G4VUserPhysicsList

G4VUserActionInitialization

Global: only one instance exists in memory, shared by all threads.

At execution

G4VUserPrimaryGeneratorAction

G4UserRunAction

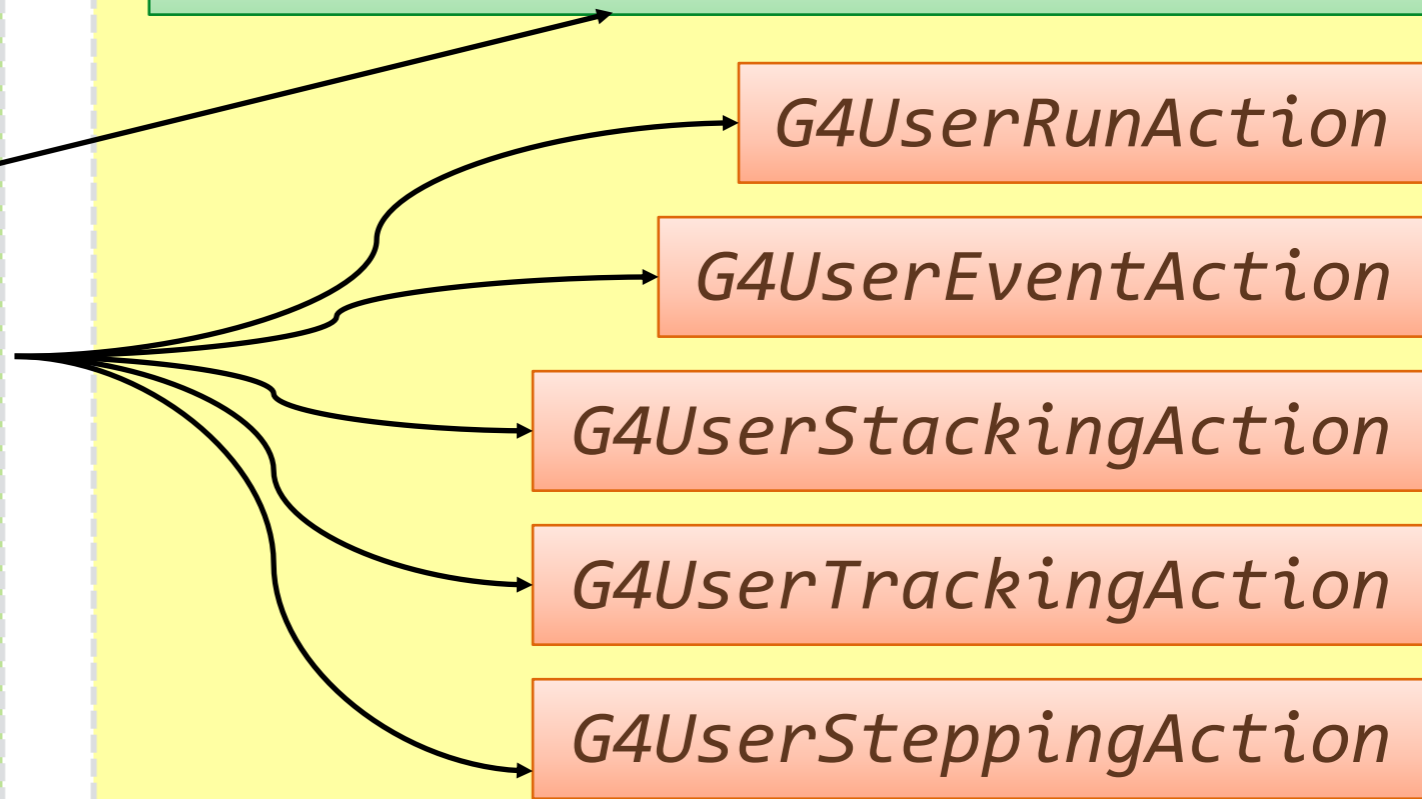
G4UserEventAction

G4UserStackingAction

G4UserTrackingAction

G4UserSteppingAction

Thread-local: an instance of each action class exists for each thread.



Note: Geant4 basic types



- Aliases for the **primitive data types** to provide cross-platform compatibility:
 - * **G4double, G4float, G4int, G4bool, G4long**
- Enhanced version of string called **G4String**
 - * **inherits from std::string ⇒ all methods and operators**
 - * **several additional methods**
- **G4ThreeVector** is a three-component class corresponding to a real physics vector (example later)

G4ThreeVector dimensions {1.0, 2.0, 3.0 };

Please, use these types for best compatibility (e.g. **G4int** instead of **int**, etc., **G4ThreeVector** when it makes sense etc.)



Part I: Units and Materials

- *System of units & constants*
 - *Definition of elements*
 - *Materials and mixtures*
 - *NIST database*

Units in Geant4



- Don't use default units!

- * When specifying dimensions, always multiply by an appropriate unit:

```
G4double width = 12.5 * m;  
G4double density = 2.7 * g/cm3;
```

- * Most common units are defined in **CLHEP** library (included in Geant4):

```
▶ G4SystemOfUnits.hh
```

```
▶ CLHEP/SystemOfUnits.hh
```

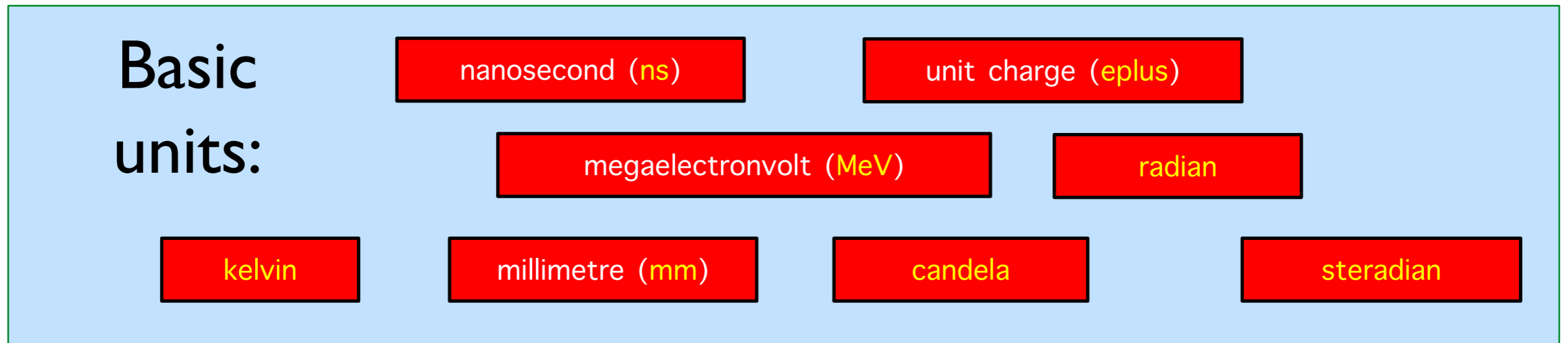
- * You can define new units (not shown here)

- Output data in terms of a specific unit:

- * divide a value by the unit

```
G4cout << dE / MeV << " (MeV)" << G4endl;
```

System of units



- All other units derived from the basic ones.
- **Useful feature:** Geant4 can select the most appropriate unit to use
 - * specify the *category* for the data (Length, Time, Energy, etc...):

```
G4cout << G4BestUnit(StepSize, "Length");
```

- **StepSize** will be printed in km, m, mm or ... fermi, depending on its actual value

- Different levels of material description:
 - * isotopes → **G4Isotope**
 - * elements → **G4Element**
 - * molecules, compounds and mixtures → **G4Material**

- Attributes associated:
 - * temperature, pressure, state, density

- **G4Isotope** and **G4Element** describe properties of the atoms:
 - * Atomic number, number of nucleons, mass of a mole, shell energies, cross-sections per atoms, etc...

- **G4Material** describes the **macroscopic** properties of the matter:
 - * Temperature, pressure, state, density
 - * Radiation length, absorption length, etc...

- **G4Material** is used by tracking, geometry and physics

Making elements

- Isotopes can be assembled into elements

```
G4Isotope (const G4String& name,  
          G4int      z,      // atomic number  
          G4int      n,      // number of nucleons  
          G4double   a ); // mass of mole
```

Not number of neutrons! →

← Do not forget unit (g/mole)

- ... building elements as follows:

```
G4Element (const G4String& name,  
          const G4String& symbol, // element symbol  
          G4int      nIso ); // n. of isotopes  
G4Element::AddIsotope (G4Isotope* iso, // isotope  
                      G4double relAbund); // fraction of atom per volume
```

- Otherwise, create G4Element with natural isotopic abundance:

```
G4Element (const G4String& name,  
          const G4String& symbol,  
          G4int      z,      // atomic number  
          G4double   a ); // mass of mole
```

← Do not forget unit (g/mole)

Elements and compounds

□ Single-element material

```
G4double z, a, density;  
density = 1.390*g/cm3;  
a = 39.95*g/mole;  
G4Material* lAr = new G4Material("liquidAr", z=18, a, density);
```

□ Molecule material (composition by number of atoms):

```
a = 1.01*g/mole;  
G4Element* e1H = new G4Element("Hydrogen", symbol="H", z=1., a);  
  
a = 16.00*g/mole;  
G4Element* e1O = new G4Element("Oxygen", symbol="O", z=8., a);  
  
density = 1.000*g/cm3;  
G4Material* H2O = new G4Material("Water", density, ncomponents=2);  
H2O->AddElement(e1H, natoms=2);  
H2O->AddElement(e1O, natoms=1);
```

□ Composition by fraction of mass

```
a = 14.01*g/mole;  
G4Element* e1N = new G4Element(name="Nitrogen", symbol="N", z= 7., a);  
a = 16.00*g/mole;  
G4Element* e1O = new G4Element(name="Oxygen", symbol="O", z= 8., a);  
density = 1.290*mg/cm3;  
G4Material* Air = new G4Material(name="Air", density, ncomponents=2);  
Air->AddElement(e1N, 70.0*perCent);  
Air->AddElement(e1O, 30.0*perCent);
```

□ Composition of mixtures

```
G4Element* e1C = ...; // define "carbon" element  
G4Material* SiO2 = ...; // define "quartz" material  
G4Material* H2O = ...; // define "water" material  
density = 0.200*g/cm3;  
  
G4Material* aerogel = new G4Material("Aerogel",  
                                     density, ncomponents=3);  
aerogel->AddMaterial(SiO2, fractionmass=62.5*perCent);  
aerogel->AddMaterial(H2O, fractionmass=37.4*perCent);  
aerogel->AddElement(e1C, fractionmass= 0.1*perCent);
```

An example: a gas

- Necessary to specify temperature and pressure

* affect dE/dx calculations, thermal scattering

```
G4double density = 27. * mg/cm3;  
G4double temperature = 325. * kelvin;  
G4double pressure = 50. * atmosphere;  
  
G4Material* CO2 = new G4Material("CO2Gas", density,  
    ncomponents=2, kStateGas, temperature, pressure);  
CO2->AddElement(C, natoms = 1);  
CO2->AddElement(O, natoms = 2);
```

- Absolute vacuum does not exist: gas at very low ρ !

* Cannot define materials with $\rho=0$

```
G4double rho = 1.e-25*g/cm3;  
G4double pr = 3.e-18*pascal;  
G4Material* Vacuum = new G4Material("interGalactic", Z, A, rho,  
kStateGas, temperature, pr);
```

NIST material database



- No need to predefine elements and materials
- Retrieve elements and materials from NIST manager:

```
G4NistManager* manager = G4NistManager::Instance();  
G4Material* H2O = manager->FindOrBuildMaterial("G4_WATER");  
G4Material* air = manager->FindOrBuildMaterial("G4_AIR");  
G4Material* vacuum = manager->FindOrBuildMaterial("G4_Galactic");  
G4Element* Si = manager->FindOrBuildElement("Si");
```

- UI commands:

`/material/nist/printElement`

← print defined elements

`/material/nist/listMaterials`

← print defined materials

▶ `G4NistManager.hh`

NIST material database



- NIST database for materials is imported inside Geant4
 - * <http://physics.nist.gov/PhysRefData>
- UI commands specific for handling materials
- The best accuracy for the most relevant parameters guaranteed:
 - * Density
 - * Mean excitation potential
 - * Chemical bounds
 - * Element composition
 - * Isotope composition
 - * Various corrections

Z	A	m	error	(%)	A _{eff}
14	Si 22	22.03453	(22)		28.0855(3)
	23	23.02552	(21)		
	24	24.011546	(21)		
	25	25.004107	(11)		
	26	25.992330	(3)		
	27	26.98670476	(17)		
	28	27.9769265327	(20)	92.2297 (7)	
	29	28.97649472	(3)	4.6832 (5)	
	30	29.97377022	(5)	3.0872 (5)	
	31	30.97536327	(7)		
	32	31.9741481	(23)		
	33	32.978001	(17)		
	34	33.978576	(15)		
	35	34.984580	(40)		
	36	35.98669	(11)		
	37	36.99300	(13)		
	38	37.99598	(29)		
	39	39.00230	(43)		
	40	40.00580	(54)		
	41	41.01270	(64)		
	42	42.01610	(75)		

- Natural isotope compositions
- More than 3000 isotope masses

NIST materials

□ NIST elementary materials:

* H → Cf (Z = 1 → 98)

□ NIST compounds:

* e.g. “G4_ADIPOSE_TISSUE_IRCP”

□ HEP and Nuclear materials:

* e.g. Liquid Ar, PbWO

□ Possible to build mixtures of NIST and user-defined materials

```
=====  
### Elementary Materials from the NIST Data  
=====  
Z Name ChFormula density(g/cm^3) I(eV)  
=====  
1 G4_H H_2 8.3748e-05 19.2  
2 G4_He 0.000166322 41.8  
3 G4_Li 0.534 40  
4 G4_Be 1.848 63.7  
5 G4_B 2.37 76  
6 G4_C 2 81  
7 G4_N N_2 0.0011652 82  
8 G4_O O_2 0.00133151 95  
9 G4_F 0.00158029 115  
10 G4_Ne 0.000838505 137  
11 G4_Na 0.971 149
```

```
=====  
### Compound Materials from the NIST Data Base  
=====  
N Name ChFormula density(g/cm^3) I(eV)  
=====  
13 G4_Adipose_Tissue 0.92 63.2  
1 0.119477  
6 0.63724  
7 0.00797  
8 0.232333  
11 0.0005  
12 2e-05  
15 0.00016  
16 0.00073  
17 0.00119  
19 0.00032  
20 2e-05  
26 2e-05  
30 2e-05  
4 G4_Air 0.00120479 85.7  
6 0.000124  
7 0.755268  
8 0.231781  
18 0.012827  
2 G4_CsI 4.51 553.1  
53 0.47692  
55 0.52308
```



Part II: Geometry

- *Basic concepts*
- *Implementation*
- *Tools for geometry checking*

Describe your detector

- A detector geometry is made of a number of volumes
- The largest volume is called **World** volume
 - * It **must** contain **all other volumes**
- Derive your own concrete class from **G4VUserDetectorConstruction** abstract base class
- Implementing the pure virtual method **Construct ()**:
 - * Define shapes/solids required to describe the geometry
 - * Construct all necessary materials
 - * Construct and place volumes of your detector geometry
 - * (Define "sensitivity" properties associated to volumes)
 - * (Associate magnetic field to detector regions)
 - * (Define visualization attributes for the detector elements)

- Implement a class inheriting from the abstract base class

G4VUserDetectorConstruction:

```
class MyDetector : public G4VUserDetectorConstruction {
public:
    virtual G4VPhysicalVolume* Construct();           // required

    virtual void ConstructSDAndField();              // optional
    // ...
};
```

- Create an instance in the main program:

```
MyDetector* detector = new MyDetector();
runManager->SetUserInitialization(detector);
```

- Note: Split the implementation into more classes and methods! (good programming practice)

* especially for complex geometries!

- Note2: you should not delete the **MyDetector** instance! Run manager does that automatically.

- Method **Construct()**

- * Define materials
- * Define solids and volumes of the geometry
- * Build the tree hierarchy of volumes
- * Define visualization attributes
- * Return the world physical volume!



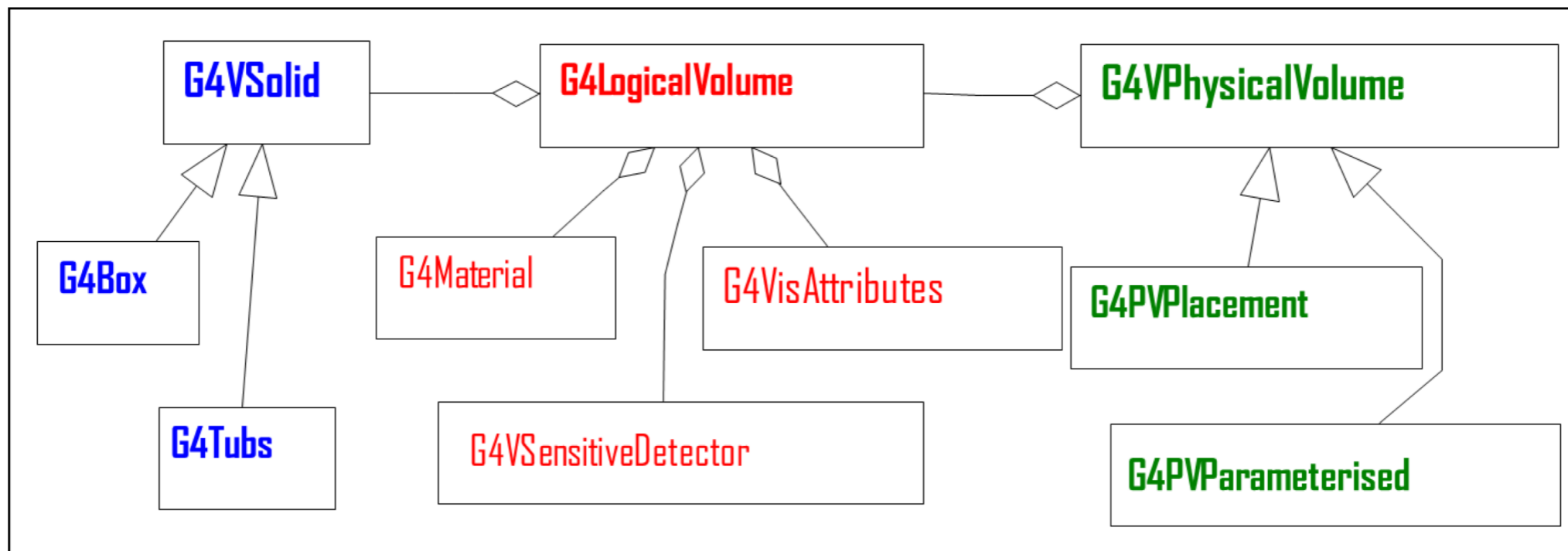
- Method **ConstructSDAndField()**

MT

- * Assign magnetic field to volumes / regions
- * Define sensitive detectors and assign them to volumes

Three conceptual layers

- **G4VSolid**
 - * *Shape, size*
- **G4LogicalVolume**
 - * *Hierarchy of volumes, material, sensitivity, magnetic field*
- **G4VPhysicalVolume**
 - * *Position, rotation. The same logical volume can be placed many times (repeated modules)*

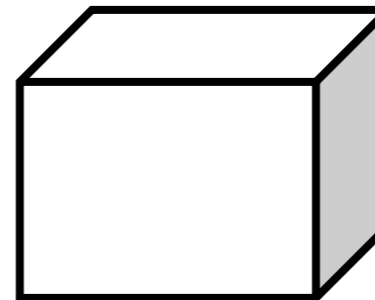


Define detector geometry

□ Basic strategy

```
G4VSolid* pBoxSolid =  
    new G4Box("aBoxSolid",  
             1.*m, 2.*m, 3.*m);
```

Solid: shape and size.



Step 1
Create the
geom. object:
box

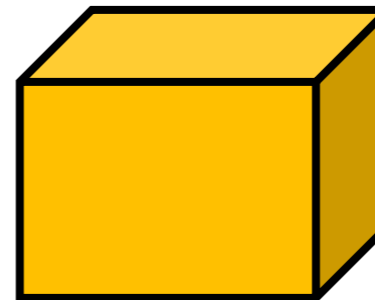
Define detector geometry

□ Basic strategy

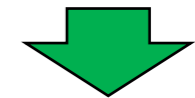
Logical volume : + material, sensitivity, etc.

```
G4VSolid* pBoxSolid =  
    new G4Box("aBoxSolid",  
            1.*m, 2.*m, 3.*m);
```

```
G4LogicalVolume* pBoxLog =  
    new G4LogicalVolume( pBoxSolid,  
                        pBoxMaterial, "aBoxLog", 0, 0, 0);
```



Step 1
Create the
geom. object:
box



Step 2
Assign properties
to object : material

Define detector geometry

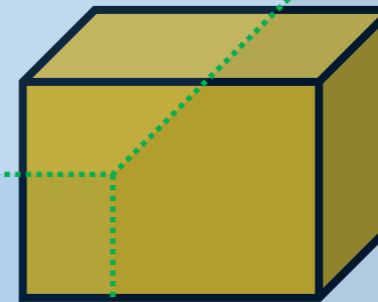
Basic strategy

Physical volume : + rotation and position

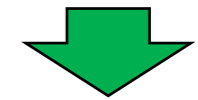
```
G4VSolid* pBoxSolid =  
new G4Box("aBoxSolid",  
1.*m, 2.*m, 3.*m);
```

```
G4LogicalVolume* pBoxLog =  
new G4LogicalVolume(pBoxSolid,  
pBoxMaterial, "aBoxLog", 0, 0, 0);
```

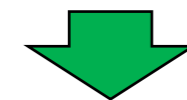
```
G4VPhysicalVolume* aBoxPhys =  
new G4PVPlacement(pRotation,  
G4ThreeVector(posX, posY, posZ),  
pBoxLog, "aBoxPhys", pMotherLog, 0, copyNo);
```



Step 1
Create the
geom. object:
box

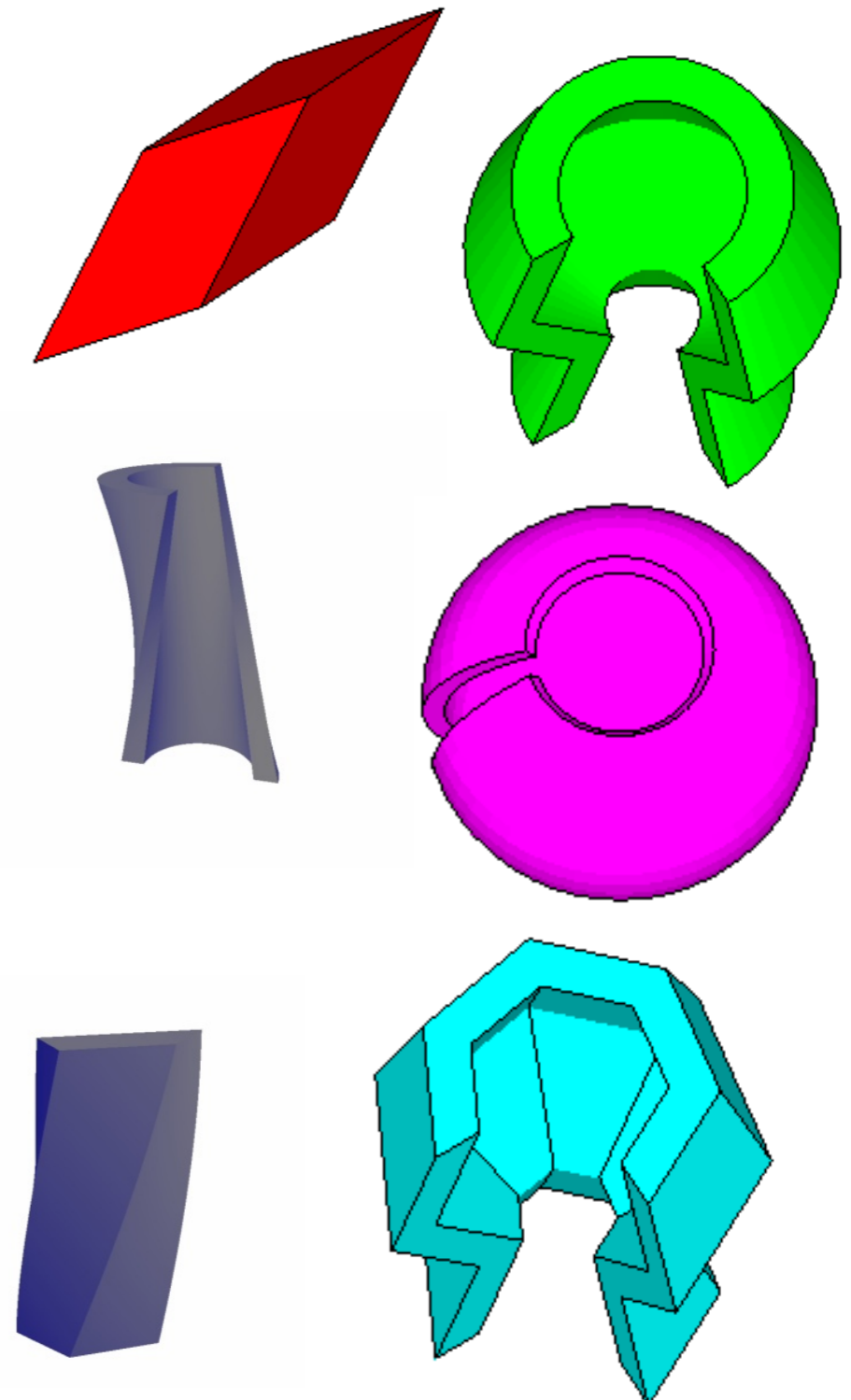


Step 2
Assign properties
to object : material



Step 3
Place it in the coordinate system of
mother volume

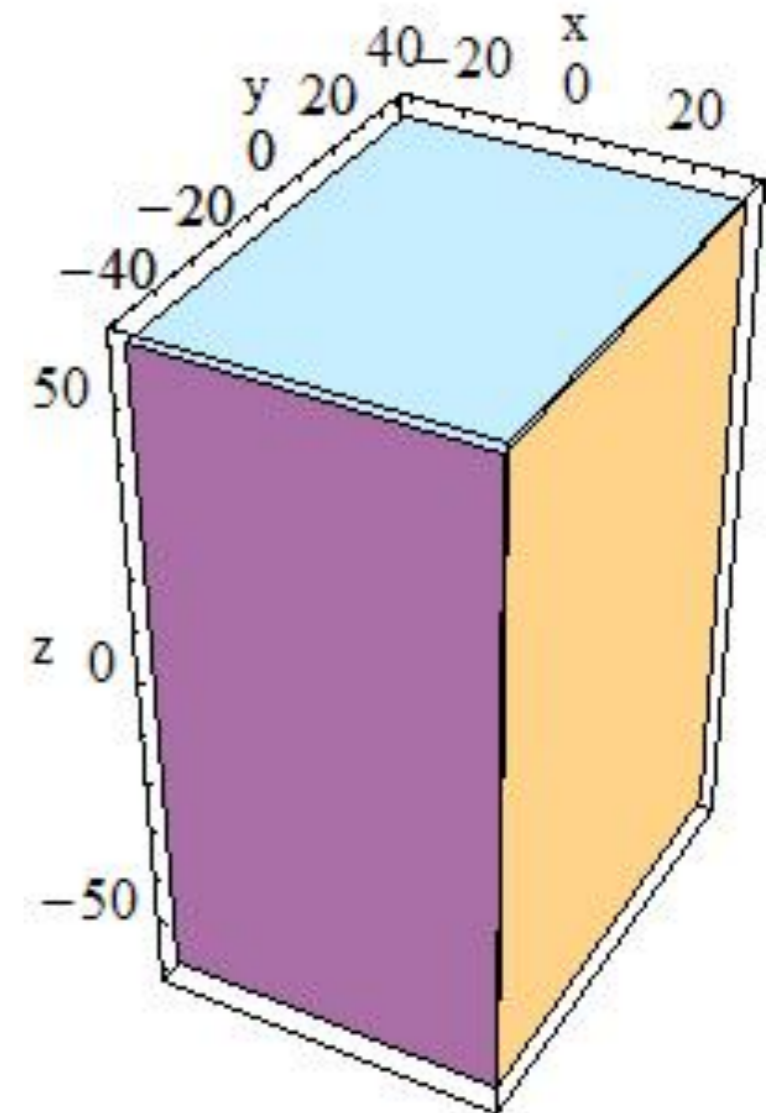
- **CSG (Constructed Solid Geometry) solids**
 - * G4Box, G4Tubs, G4Cons, G4Trd, ...
 - * Analogous to simple GEANT3 CSG solids
- **Specific solids (CSG like)**
 - * G4Polycone, G4Polyhedra, G4Hype, ...
 - * G4TwistedTubs, G4TwistedTrap, ...
- **BREP (Boundary REPresented) solids**
 - * G4BREPSolidPolycone, G4BSplineSurface, ...
 - * Any order surface
- **Boolean solids**
 - * G4UnionSolid, G4SubtractionSolid, ...



CGS: G4Box

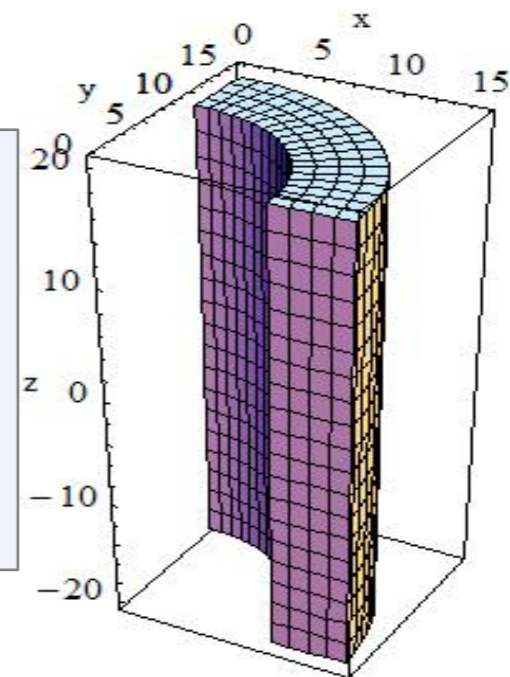
```
G4Box(const G4String& pname, // name
      G4double pX, // half-length in X
      G4double pY, // half-length in Y
      G4double pZ, // half-length in Z);
```

Note the half-length!

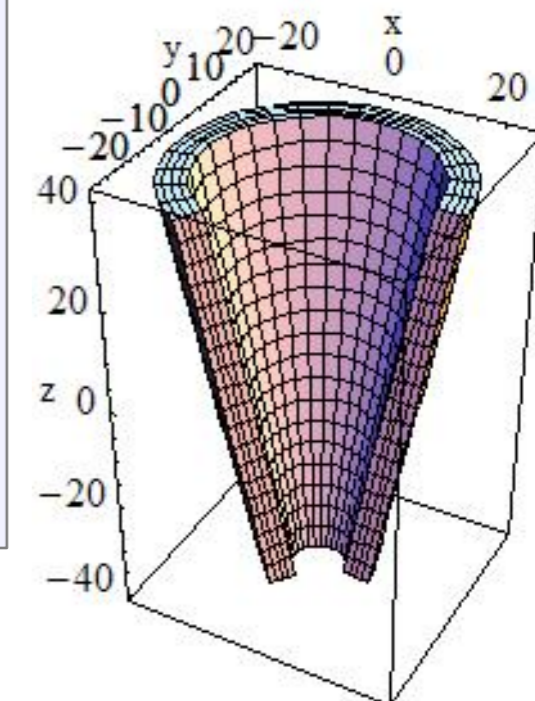


CGS: G4Tubs & G4Cones

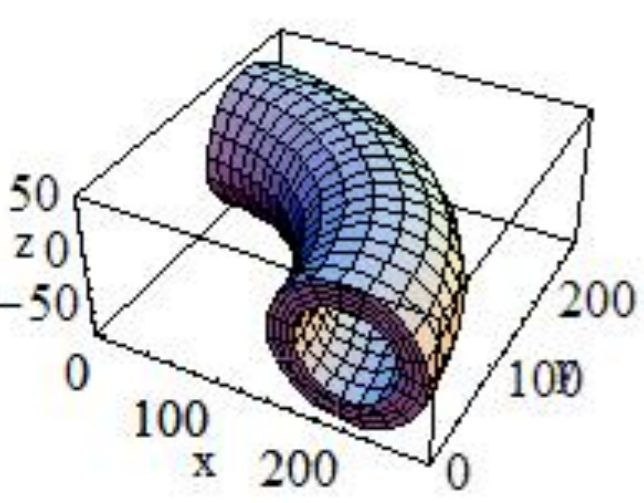
```
G4Tubs (const G4String& pname, // name
        G4double pRmin, // inner radius (0)
        G4double pRmax, // outer radius
        G4double pDz, // Z half! length
        G4double pSphi, // starting Phi (0)
        G4double pDphi); // segment angle (twopi)
```



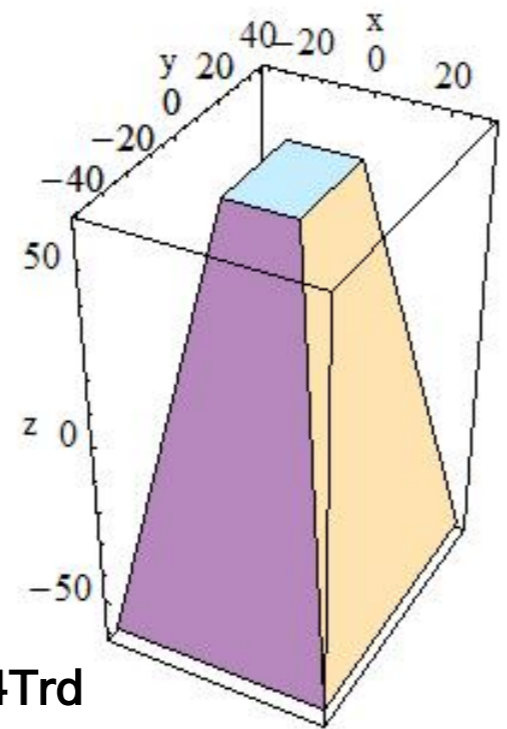
```
G4Cons (const G4String& pname, // name
        G4double pRmin1, // inner radius -pDz
        G4double pRmax1, // outer radius -pDz
        G4double pRmin2, // inner radius +pDz
        G4double pRmax2, // outer radius +pDz
        G4double pDz, // Z half length
        G4double pSphi, // starting Phi
        G4double pDphi); // segment angle
```



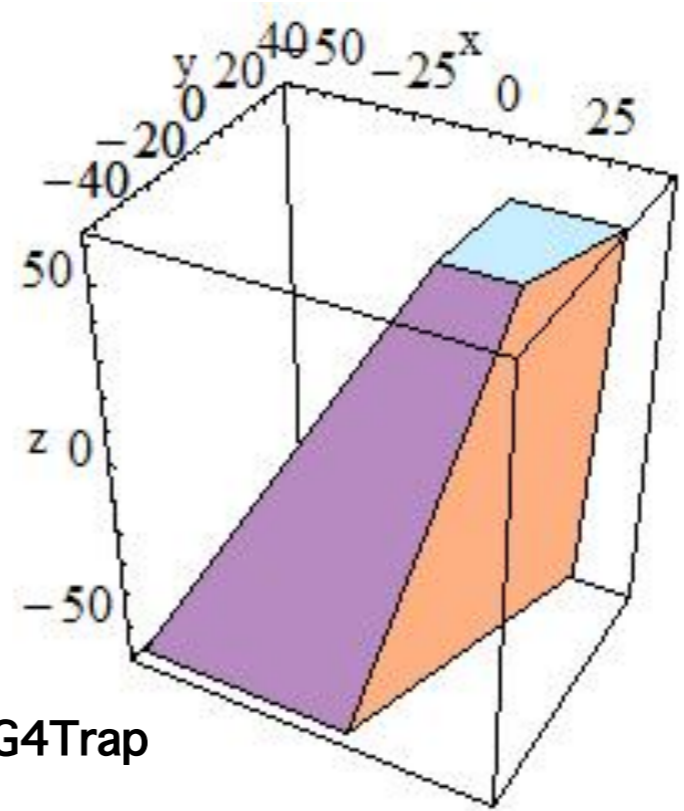
Other CGS solids



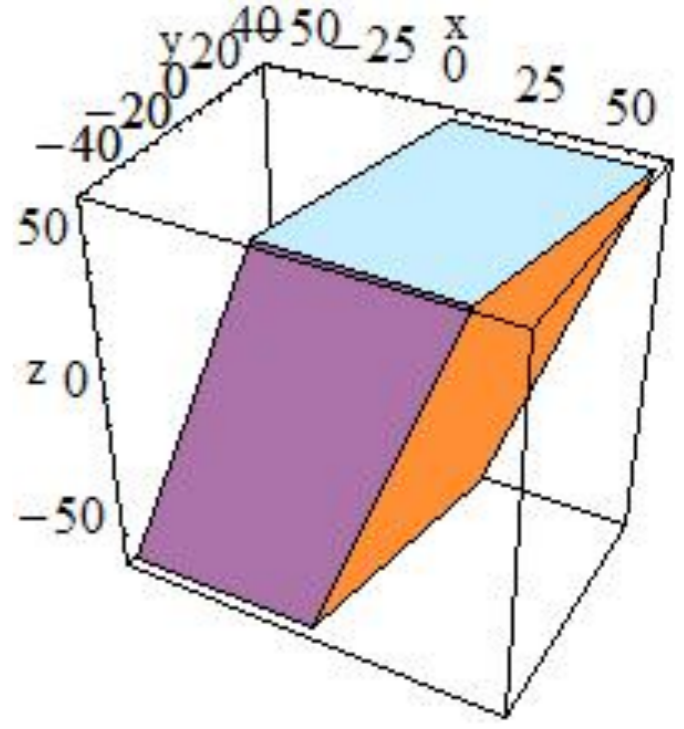
G4Torus



G4Trd

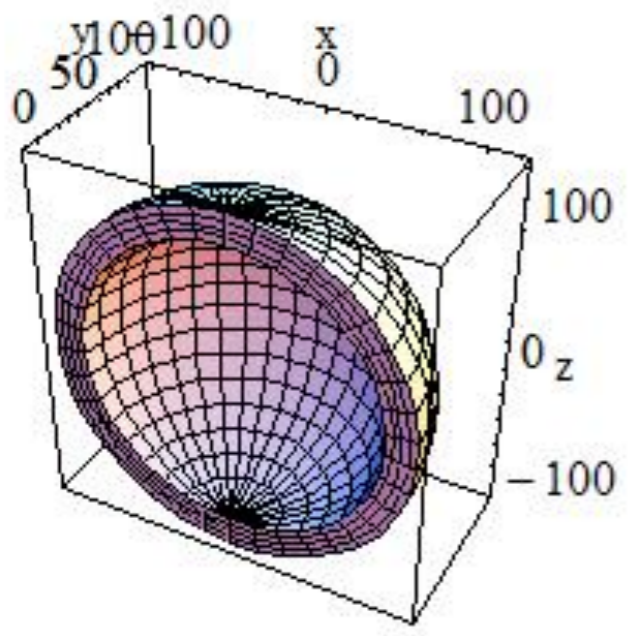


G4Trap

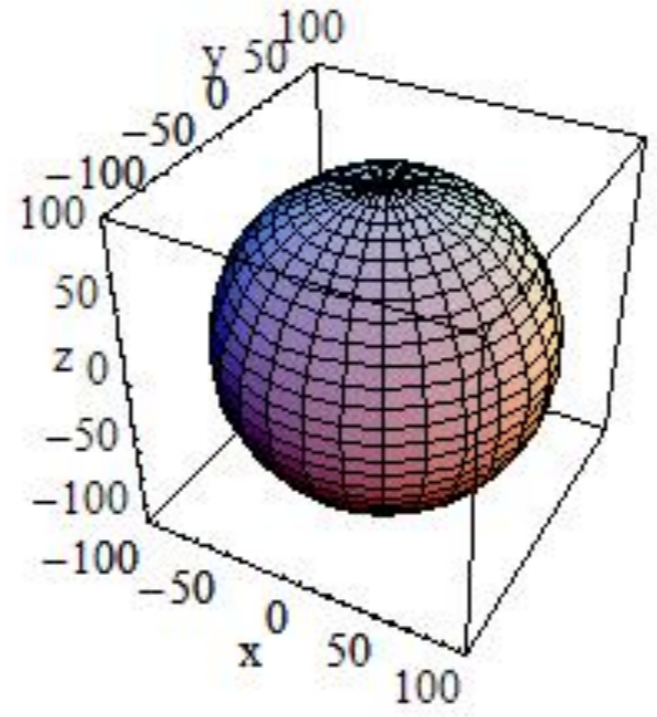


G4Para
(parallelepiped)

G4Sphere



G4Orb
(full solid
sphere)

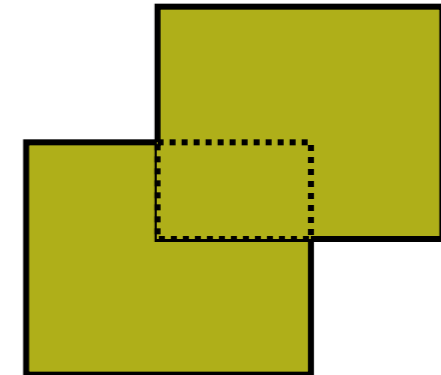


Check [Section 4.1.2](#) of Geant4 Application Developers Guide for [all available shapes](#)

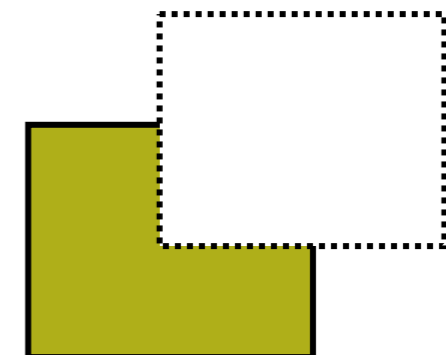
Boolean solids

- Solids can be **combined** using boolean operations:
 - * **G4UnionSolid,**
G4SubtractionSolid,
G4IntersectionSolid
 - * Requires: 2 solids, 1 boolean operation, and an (optional) transformation for the 2nd solid
 - * 2nd solid is positioned relative to the coordinate system of the 1st solid
 - * Result of boolean operation becomes a solid
→ re-usable in a boolean operation
- Solids to be combined can be either CSG or other Boolean solids
- Note: tracking **cost** for the navigation in a complex Boolean solid is proportional to the number of constituent CSG solids

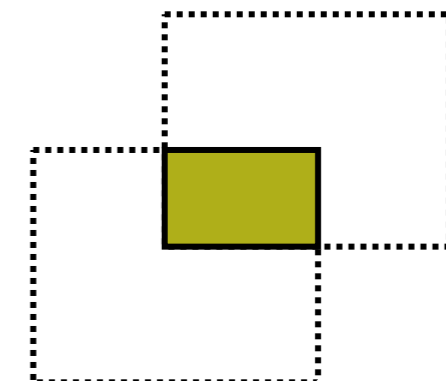
G4UnionSolid



G4SubtractionSolid



G4IntersectionSolid



Boolean solids – an example

```
G4VSolid* box = new G4Box("Box", 50*cm, 60*cm, 40*cm);
G4VSolid* cylinder =
    new G4Tubs("Cylinder", 0., 50.*cm, 50.*cm, 0., twopi);

G4VSolid* union =
    new G4UnionSolid("Box+Cylinder", box, cylinder);

G4VSolid* subtract =
    new G4SubtractionSolid("Box-Cylinder", box, cylinder,
        0, G4ThreeVector(30.*cm, 0., 0.));

G4RotationMatrix* rm = new G4RotationMatrix();
rm->RotateX(30.*deg);
G4VSolid* intersect =
    new G4IntersectionSolid("Box&&Cylinder",
        box, cylinder, rm, G4ThreeVector(0., 0., 0.));
```

Boolean solid - example



Logical volumes

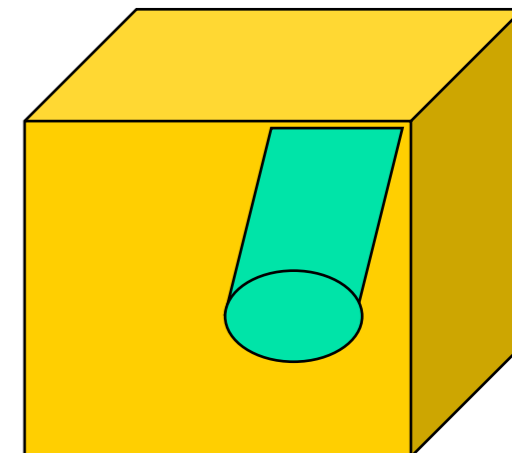
- Contains all information of volume **except position**:
 - * Shape and dimension (**G4VSolid**)
 - * Material, sensitivity, visualization attributes
 - * Position of daughter volumes
 - * Magnetic field, User limits
- Physical volumes of same type can share a logical volume.
- The pointers to solid and material must be **not nullptr**

```
G4LogicalVolume(G4VSolid* pSolid,  
                G4Material* pMaterial,  
                const G4String& name,  
                G4FieldManager* pFieldMgr=0,  
                G4VSensitiveDetector* pSDetector=0,  
                G4UserLimits* pULimits=0,  
                G4bool optimise=true);
```

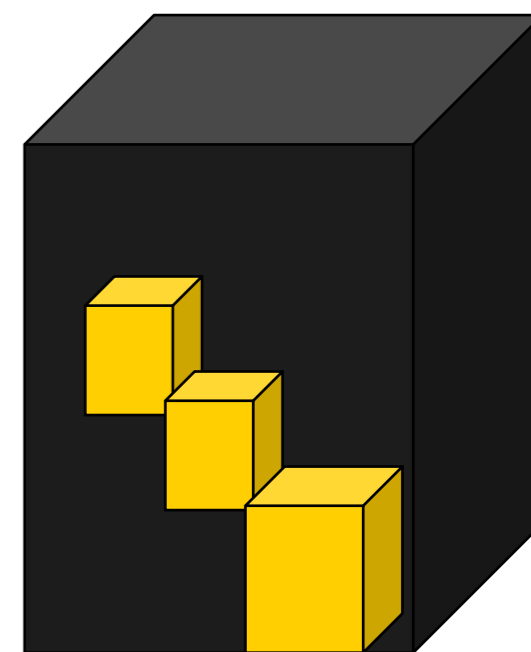
} optional

Physical volumes

- A physical volume is a **positioned instance** of a logical volume inside another logical volume (the mother volume)
- Placement (**G4PVPlacement**)
 - * it is **one** positioned volume
- Repeated: a volume placed **many times**
 - * can represent any number of volumes
 - * reduces use of memory
 - * **G4PVReplica** (= simple repetition)
 - * **G4PVParameterised** (= more complex pattern)
 - **G4PVDivision**



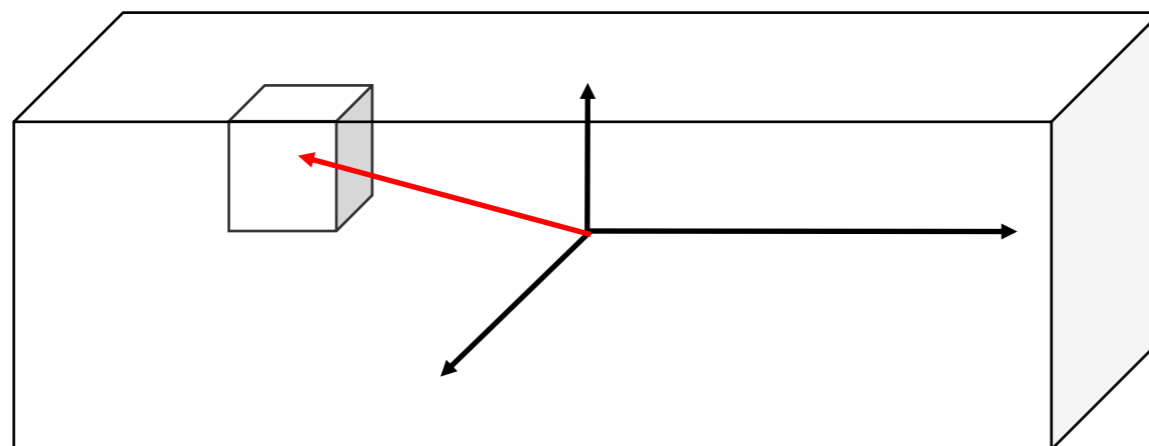
placement



repeated

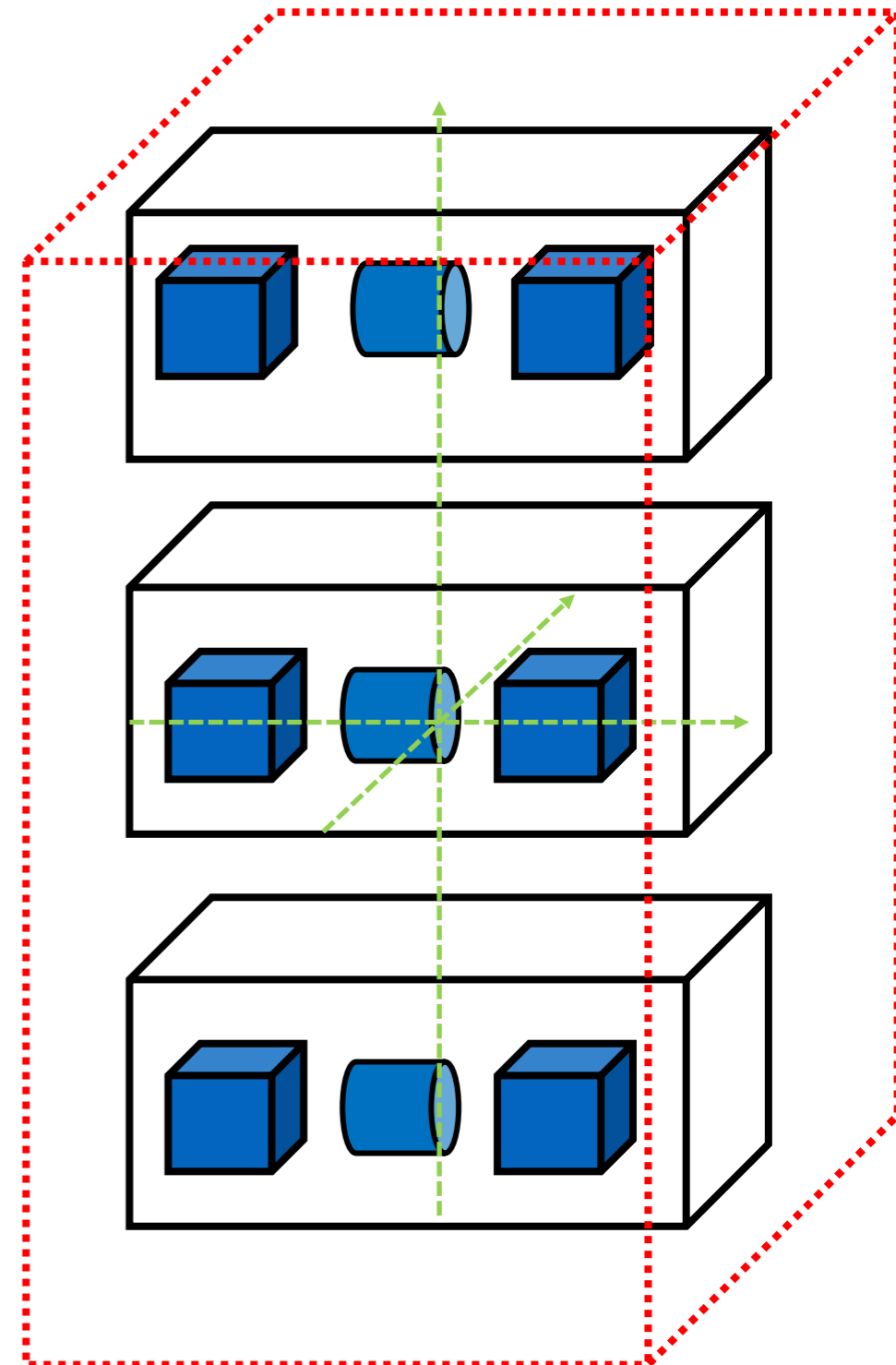
Geometry hierarchy

- A volume is placed in its mother volume
 - * Position and rotation of the daughter volume is described with respect to the **local coordinate system** of the mother volume
 - * The **origin** of the mother's local coordinate system is at the **center** of the mother volume
 - * Daughter volumes cannot protrude from the mother volume
 - * Daughter volumes cannot overlap
- The logical volume of mother knows the daughter volumes it contains
 - * It is uniquely defined to be their mother volume



Geometry hierarchy

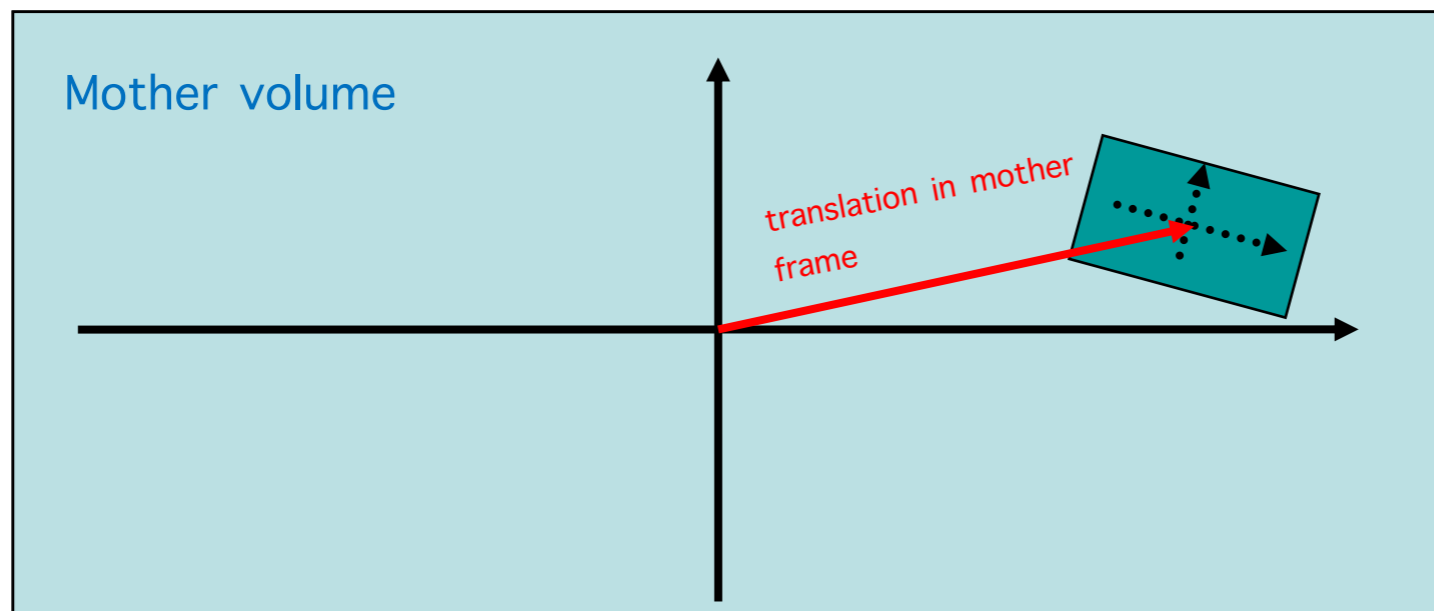
- One logical volume can be placed **more than once**. One or more volumes can be placed in a mother volume
- The mother-daughter relationship is an information of **G4LogicalVolume**
 - * If the mother volume is placed more than once, all daughters by definition appear in each placed physical volume
- The **world volume** must be a unique physical volume which fully contains all other volumes (root volume of the hierarchy)
 - * The world volume defines the **global coordinate system**. The origin of the global coordinate system is at the center of the world volume
 - * Position of a track is given with respect to the global coordinate system



- Single volume positioned **relatively** to the mother volume
 - * In a frame rotated and translated relative to the coordinate system of the **mother volume**
- A few variants available:
 - * Using **G4Transform3D** to represent the direct rotation and translation of the solid instead of the frame (*alternative constructor*)
 - * specifying the mother volume as a pointer to its physical volume instead of its logical volume
- Four constructors available
 - * **logical OR physical** volume as mother
 - * **active OR passive** transformation of the coordinate system

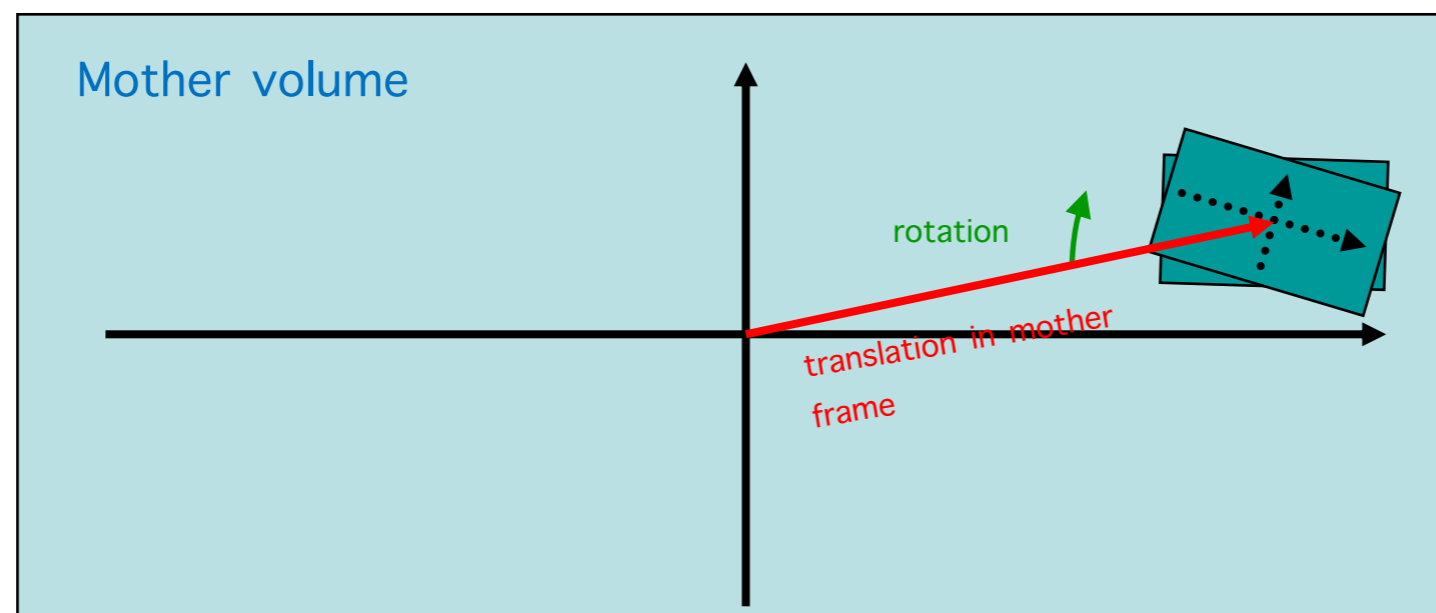
- Single volume positioned relatively to the mother volume (passive transformation)

```
G4PVPlacement(G4RotationMatrix* pRot,          // rotation of mother frame
              const G4ThreeVector& tlate,      // position in mother frame
              G4LogicalVolume* pCurrentLogical,
              const G4String& pName,
              G4LogicalVolume* pMotherLogical,
              G4bool pMany,                    // not used. Set it to false...
              G4int pCopyNo,                  // unique arbitrary index
              G4bool pSurfChk=false );        // optional overlap check
```



- Single volume positioned relatively to the mother volume (active transformation)

```
G4PVPlacement (G4Transform3D (  
    G4RotationMatrix &pRot,          // rotation in daughter frame  
    const G4ThreeVector &tlate),    // position in mother frame  
    G4LogicalVolume *pDaughterLogical,  
    const G4String &pName,  
    G4LogicalVolume *pMotherLogical,  
    G4bool pMany,                    // not used, set it to false..  
    G4int pCopyNo,                   // unique arbitrary integer  
    G4bool pSurfChk=false );        // optional overlap check
```



- Placement volume (**G4PVPlacement**): one positioned volume
 - * One physical volume represents one "real" volume
- Repeated volume: a volume placed many times
 - * One physical volume represents any number of "real" volumes
 - Reduced *use of memory*
 - Very convenient for *large voxelized* geometries
 - * Parametrized (repetitions w.r.t. copy number)
 - **G4VPVParameterisation**
 - * Replicas and Divisions (**G4PVReplica**, **G4PVDivision**)
- Notice: a repeated volume is not equivalent to a loop of placements
 - * All placements of the loop exists individually in the memory

Geometry problems

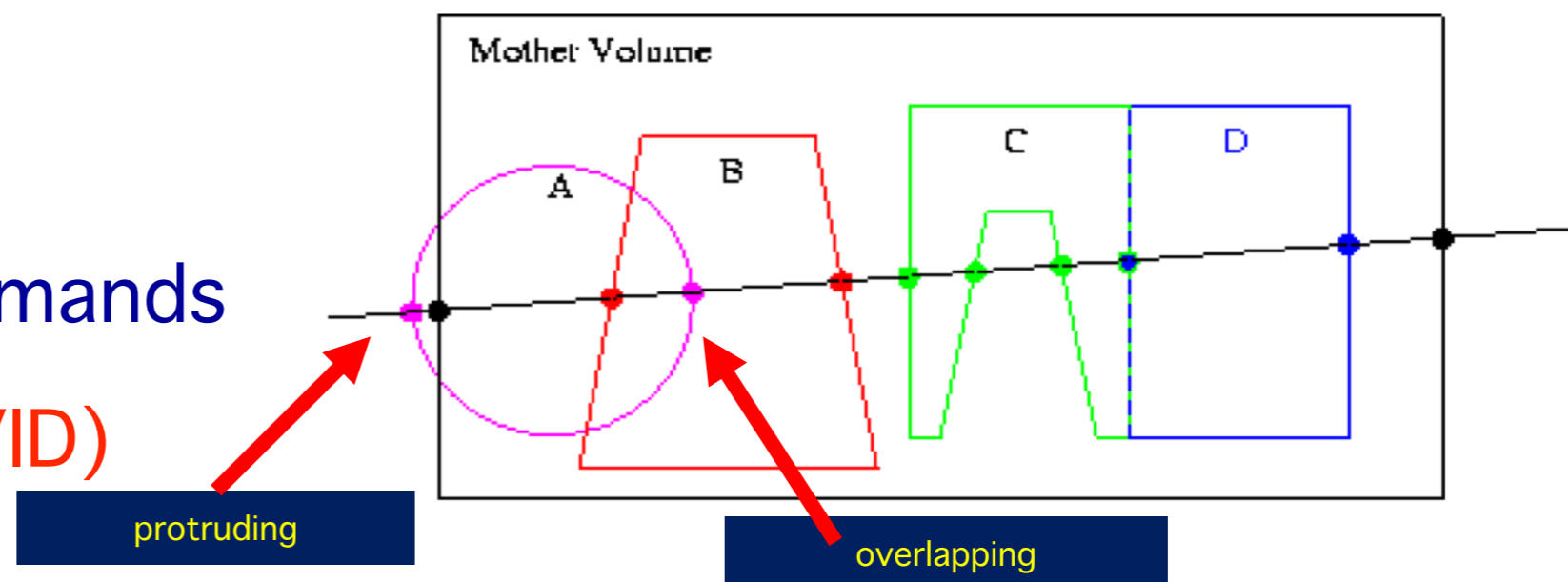
- Geant4 **does not allow** for malformed geometries, neither protruding (daughter/mother) not overlapping (sisters)
- * The behavior of navigation is **unpredictable** for such cases
- The problem of detecting overlaps between volumes is bounded by the complexity of the solid models description
- Utilities are provided for detecting wrong positioning

- * Optional checks

 - at construction

- * Kernel run-time commands

- * Graphical tools (DAVID)



Tools for geometry check



- Constructors of **G4PVPlacement** and **G4PVParameterised** have an optional argument “**pSurfChk**”

```
G4PVPlacement (G4RotationMatrix* pRot,  
               const G4ThreeVector &tlate,  
               G4LogicalVolume *pDaughterLogical,  
               const G4String &pName,  
               G4LogicalVolume *pMotherLogical,  
               G4bool pMany, G4int pCopyNo,  
               G4bool pSurfChk=false) ;
```

- * If this flag is true, **overlap check** is done at the construction
- * Some number of points are randomly sampled on the surface of creating volume
- This check requires lots of CPU time, but it is worth to **try** at least once
- Built-in run-time commands to activate verification tests for the user geometry:
 - * **/geometry/test/run** or **/geometry/test/grid_test**
 - * start **verification of geometry** for overlapping regions based on a standard grid setup, limited to the first depth level
 - * **/geometry/test/recursive_test** for all depth levels (CPU intensive!)

Tools for geometry check



```
void MyDetectorConstruction::CheckOverlaps()
{
    G4PhysicalVolumeStore* thePVStore = G4PhysicalVolumeStore::GetInstance();
    G4cout << thePVStore->size() << " physical volumes are defined" << G4endl;
    G4bool overlapFlag = false;
    G4int res = 1000;
    G4double tol = 0.; // tolerance
    for (size_t i=0; i<thePVStore->size(); i++)
    {
        overlapFlag = (*thePVStore)[i]->CheckOverlaps(res,tol,fCheckOverlapsVerbosity)
            | overlapFlag;
    }
    if (overlapFlag)
        G4cout << "Check: there are overlapping volumes" << G4endl;
}
```

Source

```
Checking overlaps for volume BeamLineSupport ... OK!
Checking overlaps for volume BeamLineCover ... OK!
Checking overlaps for volume BeamLineCover2 ... OK!
Checking overlaps for volume VacuumZone ... OK!
Checking overlaps for volume FirstScatteringFoil ... OK!
```

Output

```
.....
----- WWWWW ----- G4Exception-START ----- WWWWW -
*** G4Exception : GeomVol1002 issued by : G4PVPlacement::CheckOverlaps()
Overlap with volume already placed !
    Overlap is detected for volume BrassTube2
    with HoleNozzleSupport volume's
    local point (12.6381,12.8171,-25.1867), overlapping by at least: 3.5 mm
*** This is just a warning message ***
----- WWWWW -----
```

This method can be called at any point after run-

- A region is a **sub-set** of the geometry
- It may have its specific
 - * **Production** thresholds (**cuts**)
 - * **User** limits
 - Artificial limits affecting to the tracking, e.g. max step length, max number of steps, min kinetic energy left, etc.
 - * **Field manager**
- World logical volume is recognized as the default region. User is **not allowed** to define a region to the world logical volume

==> Task1

Link to the Tasks : <http://202.122.35.42/introduction/index.html>

Task 1 - Geometry

- Defining and using materials
- Constructing a volume using solids, logical and physical volumes
- Magnetic fields