#### NAME

**PDBFileUtil** 

#### **SYNOPSIS**

```
use PDBFileUtil ;
use PDBFileUtil qw(:all);
```

### **DESCRIPTION**

PDBFileUtil module provides the following functions:

GenerateAtomOrHetatmRecordLine, GenerateAtomRecordLine, GenerateConectRecordLine, GenerateEndRecordLine, GenerateEndRecordLine, GenerateHeaderRecordLine, GenerateHeaderRecordLine, GenerateHeaderRecordLine, GenerateHeaderRecordLine, GetAllResidues, GetChainsAndResidues, GetConectRecordLines, GetExperimentalTechnique, GetExperimentalTechnique, GetExperimentalTechniqueResolution, GetMinMaxCoords, GetPDBRecordType, GetRecordTypesCount, IsAtomRecordType, IsConectRecordType, IsEndmdlRecordType, IsHeaderRecordType, IsHeaderRecordType, IsModelRecordType, IsPDBFile, IsSeqresRecordType, IsTerRecordType, ParseAtomOrHetatmRecordLine, ParseAtomRecordLine, ParseConectRecordLine, ParseExpdtaRecordLine, ParseHeaderRecordLine, ParseHeatemRecordLine, ParseHeaderRecordLine, ParseFerRecordLine, ParseFerRecordLine, ParseFerRecordLine, ParseFerRecordLine, ParseFerRecordLine, ParseFerRecordLine, ParseFerRecordLine, ReadPDBFile

### **METHODS**

#### GenerateAtomOrHetatmRecordLine

```
$RecordLine = GenerateAtomOrHetatmRecordLine($RecordType,
    $AtomNumber, $AtomName, $AlternateLocation, $ResidueName,
    $ChainID, $ResidueNumber, $InsertionCode, $X, $Y, $Z,
    $Occupancy, $TemperatureFactor, $SegmentID,
    $ElementSymbol, $AtomCharge);
```

Returns ATOM or HETATM record line.

### GenerateAtomRecordLine

```
$RecordLine = GenerateAtomRecordLine($AtomNumber,
   $AtomName, $AlternateLocation, $ResidueName, $ChainID,
   $ResidueNumber, $InsertionCode, $X, $Y, $Z, $Occupancy,
   $TemperatureFactor, $SegmentID, $ElementSymbol, $AtomCharge);
```

Returns ATOM record line.

# GenerateConectRecordLine

```
$RecordLine = GenerateConectRecordLine($AtomNum, $BondedAtomNum1,
$BondedAtomNum2, $BondedAtomNum3, $BondedAtomNum4,
$HBondedAtomNum1, $HBondedAtomNum2, $SaltBridgedAtomNum1,
$HBondedAtomNum3, $HBondedAtomNum4, $SaltBridgedAtomNum2);
```

Returns CONECT record line.

### GenerateHeaderRecordLine

Returns HEADER record line.

## GenerateHeaderRecordTimeStamp

```
$Date = GenerateHeaderRecordTimeStamp();
```

Returns PDB header time stamp.

### GenerateHetatmRecordLine

```
$RecordLine = GenerateHetatmRecordLine($AtomNumber, $AtomName,
$AlternateLocation, $ResidueName, $ChainID, $ResidueNumber,
$InsertionCode, $X, $Y, $Z, $Occupancy, $TemperatureFactor,
$SegmentID, $ElementSymbol, $AtomCharge);
```

Returns HETATM record line.

# GenerateEndRecordLine

```
$RecordLine = GenerateEndRecordLine();
```

Returns END record line.

#### GenerateTerRecordLine

```
$RecordLine = GenerateTerRecordLine($SerialNumber, [$ResidueName,
$ChainID, $ResidueNumber, $InsertionCode]);
```

Returns TER record line.

#### GetAllResidues

```
$ResiduesDataRef = GetAllResidues($PDBRecordLinesRef);
```

Gets residue information using ATOM/HETATM records and returns a reference to a hash with following key/value pairs:

```
$ResiduesDataRef->{ResidueNames} - Array of all the residues
$ResiduesDataRef->{ResidueCount}{$ResidueName} - Count of residues
$ResiduesDataRef->{AtomResidueNames}} - Array of all ATOM residues
$ResiduesDataRef->{AtomResidueCount}{$ResidueName} - Count of
    residues in ATOM records
$ResiduesDataRef->{HetatomResidueNames} - List of all HETATM
    residues
$ResiduesDataRef->{HetatmResidueCount}{$ResidueName} - Count of
    residues HETATM records
```

ATOM/HETATM records after the first ENDMDL records are simply ingnored.

### GetChainsAndResidues

```
$ChainsDataRef = GetChainsAndResidues($PDBRecordLinesRef,
  [$RecordsSource, $GetChainResiduesBeyondTERFlag,
  $GetRecordLinesFlag]);
```

Gets chains and residue information using ATOM/HETATM or SEQRES records and returns a reference to a hash with these keys:

```
$ChainsDataRef->{ChainIDs} - List of chain IDs with 'None' for
no IDs
$ChainsDataRef->{Residues}{$ChainID} - List of residues in order
of their appearance in a chain
$ChainsDataRef->{ResidueCount}{$ChainID}{$ResidueName} - Count of
residues in a chain
```

Chains and residue data can be extacted using either ATOM/HETATM records or SEQRES records. ATOM/HETATM records after the first ENDMDL records are simply ingnored.

### GetConectRecordLines

```
$ConectRecordLinesRef = GetConectRecordLines($PDBRecordLinesRef,
$AtomNumbersMapRef);
```

Collects CONECT record lines for specific atom number, modified specified data to exclude any atom number not present in the list of specified atom numbers and returns a reference to list of CONECT record lines.

# GetExperimentalTechnique

```
$ExperimentalTechnique = GetExperimentalTechnique($PDBRecordLinesRef);
```

Returns ExperimentalTechnique value retrieved from EXPDATA record line.

### GetExperimentalTechniqueResolution

Returns Resolution and ResolutionUnits values from REMARK 2 RESOLUTION record line.

### GetMinMaxCoords

```
($XMin, $YMin, $ZMin, $XMax, $YMax, $ZMax) =
GetMinMaxCoords($PDBRecordLinesRef);
```

Returns minimum and maximum XYZ coordinates for ATOM/HETATM records.

## GetPDBRecordType

```
$RecordType = GetPDBRecordType($RecordLine);
```

Returns type of RecordLine.

## GetRecordTypesCount

\$RecordTypeDataRef = GetRecordTypesCount(\$PDBRecordLinesRef,

```
[$SpecifiedRecordType, $GetRecordLinesFlag]);
      Counts the number of each record type or a $SpecifiedRecordType and returns a reference to data type with following
      key/value pairs:
          $RecordTypeDataRef->{RecordTypes} - An array of unique record types
             in order of their presence in the file
          $RecordTypeDataRef->{Lines}{$RecordType} - Optional lines data for a
            specific record type.
IsAtomRecordType
          $Status = IsAtomRecordType($RecordLine);
      Returns 1 or 0 based on whether it's a ATOM record line.
IsConectRecordType
          $Status = IsAtomConectType($RecordLine);
      Returns 1 or 0 based on whether it's a CONECT record line.
IsEndmdIRecordType
          $Status = IsEndmdlRecordType($RecordLine);
      Returns 1 or 0 based on whether it's a ENDMDL a record line.
IsHeaderRecordType
          $Status = IsHeaderRecordType($RecordLine);
      Returns 1 or 0 based on whether it's a HEADER a record line.
IsHetatmRecordType
          $Status = IsHetatmRecordType($RecordLine);
      Returns 1 or 0 based on whether it's a HETATM a record line.
IsMasterRecordType
          $Status = IsMasterRecordType($RecordLine);
      Returns 1 or 0 based on whether it's a MASTER a record line.
IsModelRecordType
          $Status = IsModelRecordType($RecordLine);
      Returns 1 or 0 based on whether it's a MODEL record line.
IsPDBFile
          $Status = IsPDBFile($PDBFile);
      Returns 1 or 0 based on whether it's a PDB file.
IsSegresRecordType
          $Status = IsSeqresRecordType($RecordLine);
      Returns 1 or 0 based on whether it's SEQRES a record line.
IsTerRecordType
          $Status = IsTerRecordType($RecordLine);
      Returns 1 or 0 based on whether it's a TER record line.
```

Parses ATOM or HETATM record line.

ParseAtomRecordLine

ParseAtomOrHetatmRecordLine

(\$AtomNumber, \$AtomName, \$AlternateLocation, \$ResidueName, \$ChainID,

\$ResidueNumber, \$InsertionCode, \$X, \$Y, \$Z, \$Occupancy, \$TemperatureFactor, \$SegmentID, \$ElementSymbol, \$AtomCharge) =

ParseAtomOrHetatmRecordLine(\$RecordLine);

```
($AtomNumber, $AtomName, $AlternateLocation, $ResidueName, $ChainID,
            $ResidueNumber, $InsertionCode, $X, $Y, $Z, $Occupancy,
            $TemperatureFactor, $SegmentID, $ElementSymbol, $AtomCharge) =
            ParseAtomRecordLine($RecordLine);
      Parses ATOM record line.
ParseConectRecordLine
          ($AtomNum, $BondedAtomNum1, $BondedAtomNum2, $BondedAtomNum3,
             $BondedAtomNum4, $HBondedAtomNum1, $HBondedAtomNum2,
              $SaltBridgedAtomNum1, $HBondedAtomNum3, $HBondedAtomNum4,
             $SaltBridgedAtomNum2) = ParseConectRecordLine($RecordLine);
      Parses CONECT record line.
ParseExpdtaRecordLine
          ($ContinuationNum, $ExperimentalTechnique) = ParseExpdtaRecordLine($Line);
      Parses EXPDTA record line.
ParseHeaderRecordLine
          ($Classification, $DepositionDate, $IDCode) = ParseHeaderRecordLine($RecordLine);
      Parses HEADER record line
ParseHetatmRecordLine
          ($AtomNumber, $AtomName, $AlternateLocation, $ResidueName, $ChainID,
            $ResidueNumber, $InsertionCode, $X, $Y, $Z, $Occupancy,
            $TemperatureFactor, $SegmentID, $ElementSymbol, $AtomCharge) =
            ParseHetatmRecordLine($RecordLine);
      Parses HETATM record line.
ParseMasterRecordLine
           ($NumOfRemarkRecords, $NumOfHetRecords, $NumOfHelixRecords,
            $NumOfSheetRecords, $NumOfTurnRecords, $NumOfSiteRecords,
            $NumOfTransformationsRecords, $NumOfAtomAndHetatmRecords,
            $NumOfTerRecords, $NumOfConectRecords, $NumOfSeqresRecords) =
            ParseMasterRecordLine($RecordLine);
      Parses MASTER ecord line.
ParseRemark2ResolutionRecordLine
          ($Resolution, $ResolutionUnits) = ParseRemark2ResolutionRecordLine(
                                             $RecordLine);
      Parses REMARK 2 RESOLUTION record line.
ParseSegresRecordLine
          ($RecordSerialNumber, $ChainID, $NumOfResidues, $ResidueNames) =
            ParseSegresRecordLine($RecordLine);
      Parses SEQRES record line.
ParseTerRecordLine
          ($SerialNumber, $ResidueName, $ChainID, $ResidueNumber, $InsertionCode) =
            ParseTerRecordLine($RecordLine);
      Parses TER record line.
ReadPDBFile
          $PDBRecordLinesRef = ReadPDBFile($PDBFile);
      Reads PDB file and returns reference to record lines.
```

**AUTHOR** 

Manish Sud <msud@san.rr.com>

# SEE ALSO

 $File Util.pm,\ Sequence File Util.pm,\ Text Util.pm$ 

# COPYRIGHT

Copyright (C) 2018 Manish Sud. All rights reserved.

This file is part of MayaChemTools.

MayaChemTools is free software; you can redistribute it and/or modify it under the terms of the GNU Lesser General Public License as published by the Free Software Foundation; either version 3 of the License, or (at your option) any later version.