Pse-in-One: a web server for generating various modes of pseudo components of DNA, RNA, and protein sequences

Manual of stand-alone program of Pse-in-One

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Home-page: http://bioinformatics.hitsz.edu.cn/Pse-in-One/







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1. Introduction of Pse-in-One

The **Pse-in-One** web server is able to generate totally 28 different modes of pseudo components for DNA, RNA, and protein sequences, including 14 modes for DNA sequences (**Table 1**), 6 modes for RNA sequences (**Table 2**), and 8 modes for protein sequences (**Table 3**).

To the best of our knowledge, **Pse-in-One** is so far the first web server that can generate all the possible pseudo components for DNA, RNA, and protein sequences, and even those defined by users themselves, and hence it is extremely flexible.

In order to handle large dataset, the stand-alone program of **Pse-in-One** is given, which is more powerful than the Pse-in-One web server, and will be introduced in the following parts of this manual.

2. Installation

The **Pse-in-One** package can be run on Linux, Mac, and Windows systems. Download the package from http://bioinformatics.hitsz.edu.cn/Pse-in-One/download and extract it to a directory, for example, "~/usr".

To execute the **Pse-in-One** in command line environment, navigate to the "~/usr/Pse-in-One-1.0/Pse-in-One" directory and you will find three python scripts, namely "kmer.py", "acc.py" and "pse.py". The "kmer.py" is used for calculating the modes in the category nucleic acid composition or amino acid composition; The "acc.py" is used for calculating the modes in autocorrelation category. The "pse.py" is used for calculating the modes in the category pseudo nucleotide composition or pseudo amino acid composition.

3. Input/Output formats

3.1. Input format

The input file should be a valid FASTA format that consists of a single initial line beginning with a greater-than symbol (">") in the first column, followed by lines of sequence data. The words right after the ">" symbol in the single initial line are optional and only used for the purpose of identification and description.

3.2. Output format

The output file formats support three choices that are suitable for downstream computational analyses, such as machine learning. The first and the default choice is the tab format. In this format, all data is separated by TABs. The second one is the LIBSVM's sparse data format. For this format, each line contains an instance and is ended by a '\n' character, like <label> <index1>:<value1> <index2>:<value2> The <label> is a category label of the sequence. The pair <index>:<value> gives a feature (attribute) value: <index> is an integer starting from 1 and <value> is a real number. The third output format is the csv format. This format is similar to the tab format. The only difference is the separation characters between data are commas.

3.3. Physicochemical Properties Selection

The Physicochemical Properties Selection file is a text file that contains a list of property names used for generating the modes in categories: autocorrelation, pseudo nucleotide composition/ pseudo amino acid composition. For example, if you want to use the "Rise", "Tilt" and "Shift" of DNA dinucleotide for calculating, the Physicochemical Properties Selection file should be written as follows:

| Rise | | | |
|-------|--|--|--|
| Tilt | | | |
| Shift | | | |

After saving this file as "propChosen.txt" and specifying it using the command "-i propChosen.txt", or just "I propChosen.txt", the above three properties will be used in calculations. Meanwhile, you can also use the command "-a True" to select all the built-in physicochemical properties for the corresponding sequence type, which can be selected by using parameter DNA, RNA or PROTEIN.

The complete lists of physicochemical properties for DNA, RNA and protein sequences used in the stand-alone program are provided in **Table 4-8**.

3.4. User-defined Physicochemical Properties

In the user-defined physicochemical index files, each index should be represented in three lines. The first line must start with a greater-than symbol (">") in the first column. The words right after the ">" symbol in the single initial line are optional and only used for the purpose of identification and description of the index. The second line lists the names of the sequence compositions (i.e. amino acids, nucleotides, dinucleotides, or trinucleotides, etc), which should be sorted in the alphabet order, such as 'A' 'C' ... 'AA' 'AC'. All the elements in this line should be separated by TAB. The corresponding values of these sequence compositions are listed in the third line, which are separated by TAB.

For example, if you defined a physicochemical property "user_property", the user-defined physicochemical index file should be written as follows,

After saving this file as "user_defined.txt" and specifying it using the command "-e user_defined.txt", or just "E user_defined.txt", the properties defined by user will be used in calculations.

4. Commands

4.1 Command line parameters for kmer.py

| | 1 0 |
|--|---------------------------------|
| Options | Interpretations |
| inputfile | The input file in FASTA format. |
| outputfile The output file stored results. | |
| k | The k value of kmer. |

| {DNA, RNA, PROTEIN} | The sequence type. |
|---------------------|--|
| -h,help | show this help message and exit. |
| -r {1,0} | Whether consider the reverse complement or not. |
| | 1 means True, 0 means False. (default = 0) |
| -f {tab, svm, csv} | The output format. (default = tab) |
| | tab Simple format, delimited by TAB. |
| | svm The LIBSVM training data format. |
| | csv The format that can be loaded into a spreadsheet |
| | program. |

4.2 Command line parameters for acc.py

| Options | Interpretations |
|---------------------|---|
| inputfile | The input file, in FASTA format. |
| outputfile | The output file stored results. |
| lag | The value of lag. |
| {DNA, RNA, PROTEIN} | The sequence type. |
| method | The method name of autocorrelation. |
| -h,help | show this help message and exit. |
| -i I | The index file user chosen. |
| -e E | The user-defined index file. |
| -a {True,False} | Choose all physicochemical indices or not. (default = |
| | False) |
| -f {tab,svm,csv} | The output format ($default = tab$). |
| | tab Simple format, delimited by TAB. |
| | svm The LIBSVM format. |
| | csv The format that can be loaded into a spreadsheet |
| | program. |

4.3 Command line parameters for pse.py

| Options | Interpretations |
|---------------------|---|
| inputfile | The input file, in valid FASTA format. |
| outputfile | The outputfile stored results. |
| lamada | The value of lambda. |
| W | The value of weight. |
| {DNA, RNA, PROTEIN} | The sequence type. |
| method | The method name of pseudo components. |
| -h,help | show this help message and exit. |
| -i I | The index file user chosen. |
| -k K | The value of kmer, it works only with PseKNC method. |
| -e E | The user-defined index file, this parameter only needs to |
| | be set for PC-PseDNC-General, PC-PseTNC-General, |
| | SC-PseDNC-General, SC-PseTNC-General, PC- |
| | PseAAC-General or SC-PseAAC-General. |
| -a {True, False} | Choose all physicochemical indices or not. (default = |

| | False) |
|--------------------|--|
| -f {tab, svm, csv} | The output format ($default = tab$). |
| | tab Simple format, delimited by TAB. |
| | svm The LIBSVM format. |
| | csv The format that can be loaded into a spreadsheet |
| | program. |

4.4 Examples

For user's convenience, some examples of how to process a query sequence using command line are given below.

Example 1: Calculate the kmer composition feature vector of the query sequence and output the results in LIBSVM format.

```
kmer.py test.txt output_kmer.txt 2 DNA -f svm
```

After running the above command, the following results will be found in "output_kmer.txt" file.

0 1:0.023 2:0.034 3:0.053 4:0.023 5:0.045 6:0.086 7:0.143 8:0.06 9:0.049 10:0.15 11:0.124 12:0.049 13:0.015 14:0.064 15:0.053 16:0.03

Example 2: Calculate the auto covariance feature vector of the query sequence and output the results in LIBSVM format.

```
acc.py test.txt output_acc.txt 3 DNA TAC -a True -f svm
```

After running the above command, the following results will be found in "output_acc.txt" file.

0 1:-0.057 2:0.057 3:0.647 4:0.381 5:0.057 6:0.057 7:-0.051 8:-0.06 9:0.021 10:0.021 11:0.379 12:0.374 13:0.033 14:-0.011 15:0.413 16:0.019 17:-0.009 18:-0.009 19:-0.024 20:0.032 21:0.105 22:0.105 23:0.021 24:0.024 25:-0.008 26:-0.056 27:0.09 28:-0.088 29:-0.056 30:-0.056 31:-0.011 32:-0.008 33:-0.002 34:-0.002 35:-0.087 36:-0.085

Example 3: Calculate the PseDNC feature vector of the query sequence and output the results in CSV format.

```
pse.py test.txt output_pse.csv 3 0.2 DNA PseDNC
```

After running the above command, the following results will be found in "output_pse.csv" file.

0.01,0.016,0.024,0.01,0.021,0.04,0.066,0.028,0.023,0.069,0.057,0.023,0.007,0.02 9,0.024,0.014,0.217,0.152,0.17

Example 4: Calculate the PC-PseDNC-General feature vector of the query sequence using user-defined physicochemical index file and output the results in the CSV

format.

pse.py test.txt output_pse2.csv 3 0.2 DNA PC-PseDNC-General -e user_indices.txt -f csv

After running the above command, the following results will be found in "outut_pse2.csv" file.

0.011, 0.016, 0.025, 0.011, 0.021, 0.041, 0.068, 0.028, 0.023, 0.071, 0.059, 0.023, 0.007, 0.03, 0.025, 0.014, 0.213, 0.153, 0.161

The content of the file "test.txt" is listed as follow:

The content of the file "user_indices.txt" is listed as follow:

GGGCCGGACAATGAGAGTGTCCGCCTCC

| >user_de | fined_pro | perty | | | | | |
|----------|-----------|-------|-------|--------|--------|--------|---------|
| AA AC | AG AT | CA CC | CG CT | GA GC | GG GT | TA TC | TG TT |
| 0.063 | 1.502 | 0.783 | 1.071 | -1.376 | 0.063 | -1.664 | 0.783 - |
| 0.081 | -0.081 | 0.063 | 1.502 | -1.233 | -0.081 | -1.376 | 0.063 |

Table 1. 14 modes of DNA sequences calculated by PseDAC-General.

| Category | Mode | Description |
|-------------------|------------|---|
| Nucleic acid | Kmer | Basic kmer (1) |
| Composition | RevKmer | Reverse complementary kmer (2,3) |
| | DAC | Dinucleotide-based auto covariance (4,5) |
| | DCC | Dinucleotide-based cross covariance (4,5) |
| Autocorrelation | DACC | Dinucleotide-based auto-cross covariance (4,5) |
| Autocorrelation | TAC | Trinucleotide-based auto covariance (5) |
| | TCC | Trinucleotide-based cross covariance (5) |
| | TACC | Trinucleotide-based auto-cross covariance (5) |
| | PseDNC | Pseudo dinucleotide composition (6) |
| | PseKNC | Pseudo k-tuple nucleotide composition (7,8) |
| | PC-PseDNC- | General parallel correlation pseudo dinucleotide |
| | General | composition (9) |
| Pseudo nucleotide | PC-PseTNC- | General parallel correlation pseudo trinucleotide |
| composition | General | composition (9) |
| • | SC-PseDNC- | General series correlation pseudo dinucleotide |
| | General | composition (9) |
| | SC-PseTNC- | General series correlation pseudo trinucleotide |
| | General | composition (9) |

Table 2. 6 modes of RNA sequences calculated by PseRAC-General.

| Category | Mode | Description |
|--------------------------|--------------------|--|
| Nucleic acid composition | Kmer | Basic kmer (10) |
| Autocorrelation | DAC DCC DACC | Dinucleotide-based auto covariance (4,5,11) Dinucleotide-based cross covariance (4,5,11) Dinucleotide-based auto-cross covariance (4,5,11) |
| Pseudo nucleotide | PC-PseDNC- | General parallel correlation pseudo dinucleotide |

| composition | General SC-PseDNC- | composition (4,12) General series correlation pseudo dinucleotide |
|-------------|-----------------------|---|
| | General | composition (4,12) |

Table 3. 8 modes of protein sequences calculated by PseAAC-General.

| Category | Mode | Description |
|------------------------|-----------------------|---|
| Amino acid composition | Kmer | Basic kmer (13) |
| Autocorrelation | AC CC ACC | Auto covariance (5,11) Cross covariance (5,11) Auto-cross covariance (5,11) |
| | PC-PseAAC | Parallel correlation pseudo amino acid composition (14) |
| Pseudo amino acid | SC-PseAAC | Series correlation pseudo amino acid composition (15) |
| composition | PC-PseAAC- General | General parallel correlation pseudo amino acid composition (14,16) |
| | SC-PseAAC- General | General series correlation pseudo amino acid composition (15,16) |

Table 4. The names of the 148 physicochemical indices for dinucleotides.

| Base stacking | Protein | B-DNA twist |
|------------------------------|-------------------------------|--------------------------------|
| | induced deformability | |
| Propeller twist | Duplex stability:(freeenergy) | Duplex tability(disruptenergy) |
| Protein DNA twist | Stabilising energy of Z-DNA | Aida_BA_transition |
| Breslauer_dS | Electron_interaction | Hartman_trans_free_energy |
| Lisser_BZ_transition | Polar_interaction | SantaLucia_dG |
| Sarai_flexibility | Stability | Stacking_energy |
| Sugimoto_dS | Watson-Crick_interaction | Twist |
| Shift | Slide | Rise |
| Twist stiffness | Tilt stiffness | Shift_rise |
| Twist_shift | Enthalpy1 | Twist_twist |
| Shift2 | Tilt3 | Tilt1 |
| Slide (DNA-protein complex)1 | Tilt_shift | Twist_tilt |
| Roll_rise | Stacking energy | Stacking energy1 |
| Propeller Twist | Roll11 | Rise (DNA-protein complex) |
| Roll2 | Roll3 | Roll1 |
| Slide_slide | Enthalpy | Shift_shift |
| Flexibility_slide | Minor Groove Distance | Rise (DNA-protein complex)1 |
| Roll (DNA-protein complex)1 | Entropy | Cytosine content |
| Major Groove Distance | Twist (DNA-protein complex) | Purine (AG) content |
| Tilt_slide | Major Groove Width | Major Groove Depth |
| Free energy6 | Free energy7 | Free energy4 |
| Free energy3 | Free energy1 | Twist_roll |
| Flexibility_shift | Shift (DNA-protein complex)1 | Thymine content |
| Tip | Keto (GT) content | Roll stiffness |
| Entropy1 | Roll_slide | Slide (DNA-protein complex) |
| Twist2 | Twist5 | Twist4 |
| Tilt (DNA-protein complex)1 | Twist_slide | Minor Groove Depth |
| Persistance Length | Rise3 | Shift stiffness |
| Slide3 | Slide2 | Slide1 |
| Rise1 | Rise stiffness | Mobility to bend towards minor |
| | | • |

| | | groove |
|----------------------------|----------------------|--------------------------------|
| Dinucleotide GC Content | A-philicity | Wedge |
| DNA denaturation | Bending stiffness | Free energy5 |
| Breslauer_dG | Breslauer_dH | Shift (DNA-protein complex) |
| Helix-Coil_transition | Ivanov_BA_transition | Slide_rise |
| SantaLucia_dH | SantaLucia_dS | Minor Groove Width |
| Sugimoto_dG | Sugimoto_dH | Twist1 |
| Tilt | Roll | Twist7 |
| Clash Strength | Roll_roll | Roll (DNA-protein complex) |
| Adenine content | Direction | Probability contacting |
| | | nucleosome core |
| Roll_shift | Shift_slide | Shift1 |
| Tilt4 | Tilt2 | Free energy8 |
| Twist (DNA-protein | Tilt_rise | Free energy2 |
| complex)1 | | |
| Stacking energy2 | Stacking energy3 | Rise_rise |
| Tilt_tilt | Roll4 | Tilt_roll |
| Minor Groove Size | GC content | Inclination |
| Slide stiffness | Melting Temperature1 | Twist3 |
| Tilt (DNA-protein complex) | Guanine content | Twist6 |
| Major Groove Size | Twist_rise | Rise2 |
| Melting Temperature | Free energy | Mobility to bend towards major |
| | | groove |

Table 5. The names of the 12 physicochemical indices for trinucleotides.

| Bendability (DNAse) | Bendability (consensus) | Trinucleotide GC Content |
|------------------------|-------------------------|--------------------------|
| Consensus_roll | Consensus-Rigid | Dnase I |
| MW-Daltons | MW-kg | Nucleosome |
| Nucleosome positioning | Dnase I-Rigid | Nucleosome-Rigid |

Table 6. The names of the 6 physicochemical indices for dinucleotides.

| Twist | Tilt | Roll | |
|-------|-------|------|--|
| Shift | Slide | Rise | |

Table 7. The names of the 22 physicochemical indices for dinucleotides.

| Shift (RNA) | Hydrophilicity (RNA) |
|----------------------|-----------------------|
| Hydrophilicity (RNA) | GC content |
| Purine (AG) content | Keto (GT) content |
| Adenine content | Guanine content |
| Cytosine content | Thymine content |
| Slide (RNA) | Rise (RNA) |
| Tilt (RNA) | Roll (RNA) |
| Twist (RNA) | Stacking energy (RNA) |
| Enthalpy (RNA) | Entropy (RNA) |
| Free energy (RNA) | Free energy (RNA) |
| Enthalpy (RNA) | Entropy (RNA) |

Table 8. The names of the 547 physicochemical indices for amino acids.

| Hydrophobicity | Hydrophilicity | Mass | ANDN920101 |
|----------------|----------------|------------|------------|
| ARGP820101 | ARGP820102 | ARGP820103 | BEGF750101 |
| BEGF750102 | BEGF750103 | BHAR880101 | BIGC670101 |
| BIOV880101 | BIOV880102 | BROC820101 | BROC820102 |
| BULH740101 | BULH740102 | BUNA790101 | BUNA790102 |
| BUNA790103 | BURA740101 | BURA740102 | CHAM810101 |
| CHAM820101 | CHAM820102 | CHAM830101 | CHAM830102 |
| CHAM830103 | CHAM830104 | CHAM830105 | CHAM830106 |
| CHAM830107 | CHAM830108 | CHOC750101 | CHOC760101 |
| CHOC760102 | CHOC760103 | CHOC760104 | CHOP780101 |
| CHOP780201 | CHOP780202 | CHOP780203 | CHOP780204 |
| CHOP780205 | CHOP780206 | CHOP780207 | CHOP780208 |
| CHOP780209 | CHOP780210 | CHOP780211 | CHOP780212 |
| CHOP780213 | CHOP780214 | CHOP780215 | CHOP780216 |
| CIDH920101 | CIDH920102 | CIDH920103 | CIDH920104 |
| CIDH920105 | COHE430101 | CRAJ730101 | CRAJ730102 |
| CRAJ730103 | DAWD720101 | DAYM780101 | DAYM780201 |
| DESM900101 | DESM900102 | EISD840101 | EISD860101 |
| EISD860102 | EISD860103 | FASG760101 | FASG760102 |
| FASG760103 | FASG760104 | FASG760105 | FAUJ830101 |
| FAUJ880101 | FAUJ880102 | FAUJ880103 | FAUJ880104 |
| FAUJ880105 | FAUJ880106 | FAUJ880107 | FAUJ880108 |
| FAUJ880109 | FAUJ880110 | FAUJ880111 | FAUJ880112 |
| FAUJ880113 | FINA770101 | FINA910101 | FINA910102 |
| FINA910103 | FINA910104 | GARJ730101 | GEIM800101 |
| GEIM800102 | GEIM800103 | GEIM800104 | GEIM800105 |
| GEIM800106 | GEIM800107 | GEIM800108 | GEIM800109 |
| GEIM800110 | GEIM800111 | GOLD730101 | GOLD730102 |
| GRAR740101 | GRAR740102 | GRAR740103 | GUYH850101 |
| HOPA770101 | HOPT810101 | HUTJ700101 | HUTJ700102 |
| HUTJ700103 | ISOY800101 | ISOY800102 | ISOY800103 |
| ISOY800104 | ISOY800105 | ISOY800106 | ISOY800107 |
| ISOY800108 | JANJ780101 | JANJ780102 | JANJ780103 |
| JANJ790101 | JANJ790102 | JOND750101 | JOND750102 |
| JOND920101 | JOND920102 | JUKT750101 | JUNJ780101 |
| KANM800101 | KANM800102 | KANM800103 | KANM800104 |
| KARP850101 | KARP850102 | KARP850103 | KHAG800101 |
| KLEP840101 | KRIW710101 | KRIW790101 | KRIW790102 |
| KRIW790103 | KYTJ820101 | LAWE840101 | LEVM760101 |
| LEVM760102 | LEVM760103 | LEVM760104 | LEVM760105 |
| LEVM760106 | LEVM760107 | LEVM780101 | LEVM780102 |
| LEVM780103 | LEVM780104 | LEVM780105 | LEVM780106 |
| LEWP710101 | LIFS790101 | LIFS790102 | LIFS790103 |
| MANP780101 | MAXF760101 | MAXF760102 | MAXF760103 |
| MAXF760104 | MAXF760105 | MAXF760106 | MCMT640101 |
| MEEJ800101 | MEEJ800102 | MEEJ810101 | MEEJ810102 |
| MEIH800101 | MEIH800102 | MEIH800103 | MIYS850101 |
| • | | | |

| NAGK730101 | NAGK730102 | NAGK730103 | NAKH900101 |
|------------|------------|------------|------------|
| NAKH900102 | NAKH900103 | NAKH900104 | NAKH900105 |
| NAKH900106 | NAKH900107 | NAKH900108 | NAKH900109 |
| NAKH900110 | NAKH900111 | NAKH900112 | NAKH900113 |
| NAKH920101 | NAKH920102 | NAKH920103 | NAKH920104 |
| NAKH920105 | NAKH920106 | NAKH920107 | NAKH920108 |
| NISK800101 | NISK860101 | NOZY710101 | OOBM770101 |
| OOBM770102 | OOBM770103 | OOBM770104 | OOBM770105 |
| OOBM850101 | OOBM850102 | OOBM850103 | OOBM850104 |
| OOBM850105 | PALJ810101 | PALJ810102 | PALJ810103 |
| PALJ810104 | PALJ810105 | PALJ810106 | PALJ810107 |
| PALJ810108 | PALJ810109 | PALJ810110 | PALJ810111 |
| PALJ810112 | PALJ810113 | PALJ810114 | PALJ810115 |
| PALJ810116 | PARJ860101 | PLIV810101 | PONP800101 |
| PONP800102 | PONP800103 | PONP800104 | PONP800105 |
| PONP800106 | PONP800107 | PONP800108 | PRAM820101 |
| PRAM820102 | PRAM820103 | PRAM900101 | PRAM900102 |
| PRAM900103 | PRAM900104 | PTIO830101 | PTIO830102 |
| QIAN880101 | QIAN880102 | QIAN880103 | QIAN880104 |
| QIAN880105 | QIAN880106 | QIAN880107 | QIAN880108 |
| QIAN880109 | QIAN880110 | QIAN880111 | QIAN880112 |
| QIAN880113 | QIAN880114 | QIAN880115 | QIAN880116 |
| QIAN880117 | QIAN880118 | QIAN880119 | QIAN880120 |
| QIAN880121 | QIAN880122 | QIAN880123 | QIAN880124 |
| QIAN880125 | QIAN880126 | QIAN880127 | QIAN880128 |
| QIAN880129 | QIAN880130 | QIAN880131 | QIAN880132 |
| QIAN880133 | QIAN880134 | QIAN880135 | QIAN880136 |
| QIAN880137 | QIAN880138 | QIAN880139 | RACS770101 |
| RACS770102 | RACS770103 | RACS820101 | RACS820102 |
| RACS820103 | RACS820104 | RACS820105 | RACS820106 |
| RACS820107 | RACS820108 | RACS820109 | RACS820110 |
| RACS820111 | RACS820112 | RACS820113 | RACS820114 |
| RADA880101 | RADA880102 | RADA880103 | RADA880104 |
| RADA880105 | RADA880106 | RADA880107 | RADA880108 |
| RICJ880101 | RICJ880102 | RICJ880103 | RICJ880104 |
| RICJ880105 | RICJ880106 | RICJ880107 | RICJ880108 |
| RICJ880109 | RICJ880110 | RICJ880111 | RICJ880112 |
| RICJ880113 | RICJ880114 | RICJ880115 | RICJ880116 |
| RICJ880117 | ROBB760101 | ROBB760102 | ROBB760103 |
| ROBB760104 | ROBB760105 | ROBB760106 | ROBB760107 |
| ROBB760108 | ROBB760109 | ROBB760110 | ROBB760111 |
| ROBB760112 | ROBB760113 | ROBB790101 | ROSG850101 |
| ROSG850102 | ROSM880101 | ROSM880102 | ROSM880103 |
| SIMZ760101 | SNEP660101 | SNEP660102 | SNEP660103 |
| SNEP660104 | SUEM840101 | SUEM840102 | SWER830101 |
| TANS770101 | TANS770102 | TANS770103 | TANS770104 |
| TANS770105 | TANS770106 | TANS770107 | TANS770108 |
| TANS770109 | TANS770110 | VASM830101 | VASM830102 |
| VASM830103 | VELV850101 | VENT840101 | VHEG790101 |

| WARP780101 | WEBA780101 | WERD780101 | WERD780102 |
|---------------------------------------|------------|------------|------------|
| WERD780103 | WERD780104 | WOEC730101 | WOLR810101 |
| WOLS870101 | WOLS870102 | WOLS870103 | YUTK870101 |
| YUTK870102 | YUTK870103 | YUTK870104 | ZASB820101 |
| ZIMJ680101 | ZIMJ680102 | ZIMJ680103 | ZIMJ680104 |
| ZIMJ680105 | AURR980101 | AURR980102 | AURR980103 |
| AURR980104 | AURR980105 | AURR980106 | AURR980107 |
| AURR980108 | AURR980109 | AURR980110 | AURR980111 |
| AURR980112 | AURR980113 | AURR980114 | AURR980115 |
| AURR980116 | AURR980117 | AURR980118 | AURR980119 |
| AURR980120 | ONEK900101 | ONEK900102 | VINM940101 |
| VINM940102 | VINM940103 | VINM940104 | MUNV940101 |
| MUNV940102 | MUNV940103 | MUNV940104 | MUNV940105 |
| WIMW960101 | KIMC930101 | MONM990101 | BLAM930101 |
| PARS000101 | PARS000102 | KUMS000101 | KUMS000102 |
| KUMS000103 | KUMS000104 | TAKK010101 | FODM020101 |
| NADH010101 | NADH010102 | NADH010103 | NADH010104 |
| NADH010105 | NADH010106 | NADH010107 | MONM990201 |
| KOEP990101 | KOEP990102 | CEDJ970101 | CEDJ970102 |
| CEDJ970103 | CEDJ970104 | CEDJ970105 | FUKS010101 |
| FUKS010102 | FUKS010103 | FUKS010104 | FUKS010105 |
| FUKS010106 | FUKS010107 | FUKS010108 | FUKS010109 |
| FUKS010110 | FUKS010111 | FUKS010112 | AVBF000101 |
| AVBF000102 | AVBF000103 | AVBF000104 | AVBF000105 |
| AVBF000106 | AVBF000107 | AVBF000108 | AVBF000109 |
| YANJ020101 | MITS020101 | TSAJ990101 | TSAJ990102 |
| COSI940101 | PONP930101 | WILM950101 | WILM950102 |
| WILM950103 | WILM950104 | KUHL950101 | GUOD860101 |
| JURD980101 | BASU050101 | BASU050102 | BASU050103 |
| SUYM030101 | PUNT030101 | PUNT030102 | GEOR030101 |
| GEOR030102 | GEOR030103 | GEOR030104 | GEOR030105 |
| GEOR030106 | GEOR030107 | GEOR030108 | GEOR030109 |
| ZHOH040101 | ZHOH040102 | ZHOH040103 | BAEK050101 |
| HARY940101 | PONJ960101 | DIGM050101 | WOLR790101 |
| OLSK800101 | KIDA850101 | GUYH850102 | GUYH850103 |
| GUYH850104 | GUYH850105 | ROSM880104 | ROSM880105 |
| JACR890101 | COWR900101 | BLAS910101 | CASG920101 |
| CORJ870101 | CORJ870102 | CORJ870103 | CORJ870104 |
| CORJ870105 | CORJ870106 | CORJ870107 | CORJ870108 |
| MIYS990101 | MIYS990102 | MIYS990103 | MIYS990104 |
| MIYS990105 | ENGD860101 | FASG890101 | |
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