

OptiChamber Software (version 2.0) for Modeling Laser-Pumping Trains

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1. Overview

OptiChamber v.2.0. is intended for computational modeling of optical trains containing:

- multiple diode bars pumping unit,
- laser rod being pumped,
- and reflector surrounding laser rod.

Present software implies 2D geometry, i.e., all the above elements are assumed to have infinite length along the laser beam direction. This enables to omit consideration of the fringe effects and describe radiation transfer within the cross-section plane only. Version 2.0 of the software provides modeling of continuous (stationary) pumping for five pre-defined or user-defined configurations, indicated by the type number, $iCf_g=1\dots 5$, which containing diode bars and rods. For these configurations the modeling software package yields:

- two-dimensional distribution of the power density absorbed in the area of active medium [W/cm^3];
- histogram of area distribution for different levels of the absorbed power density [percents of the total area];
- maximum value of the absorbed power density [W/cm^3];
- total lineal (per unit length) power absorbed in the active medium [W/cm];

- efficiency, that is, the ratio of total power absorbed in the active medium to total power emitted by the lamp or diode [percents].

2. First-Step Installation

Full-operation version of the OptiChamber software is installed in two steps. First you install the Pre-Registration Copy, whose setup does not require any serial number. It allows changing only few parameters of the optical train and can serve as a demo version. The second-step installation, which requires entering a serial number (to be discussed in the next section), upgrades the demo copy to the full-operation version.

2.1. System requirements

- Computer: IBM PC Compatible.
- CPU type: 80486 or higher.
- RAM: 16Mb or more.
- Video Card: providing at least SVGA 800x600, 256 colors (16-bit Hi Color mode is recommended for adequate reproduction of two-dimensional plots).
- Available disk space: about 10 MB
- Operation system: Windows 95/98/2000.

2.2. Setup procedure

- Insert the Setup Disk in your CD ROM drive or download the setup files from Internet to hard disk.
- Run file **OC2_INSTALL.EXE** on this drive.
- Follow the instructions on your screen.
During the installation you will be prompted to enter your and your company's name and to select the destination folder in which the program files are stored (by default, C:\Program Files\OptiChamber2). When the setup procedure finishes, the following files appear in the selected folder:

Main files

- **OptiChamber.exe (shell file)**- main program file providing end-user interface,
- **OCCalc2Demo.exe (kernel file)**- program providing calculations.

Supplementary files

- **OptiChamber.rtf** – Short User's Guide,
- **Readme.txt** - list of last-minute changes,
- **pump.dan** - input data file for OCCalc2,
- **pump1.dat** - output data file of OCCalc2, which is used for representation of final results,
- **pump2.dat** - auxiliary file containing additional information output of OCCalc2,
- files representing sample lamp (*.ls) and laser rod (*.rs) emission and absorption spectra,
- *.shp files representing sample reflector's shapes.

File **pump.ini** - default OptiChamber configuration - appear in the OptiChamber folder after the first run and saving configuration.

The following extensions are used to by default to identify additional files that can be read or written by the software from/to the OptiChamber home or another directory indicated by user:

- *.dan - OptiChamber configuration file (parameters to be inserted into user interface fields of OptiChamber),
- *.dat - calculation results in OptiChamber native or Surfer format,
- *.prn - two-dimensional distribution of the absorbed power density in MathCad format,
- *.ls - lamp or diode emission spectra,
- *.rs - laser rod absorption spectrum,
- *.shp - file describing custom-shape reflector.

2.3. Uninstall procedure

You can readily remove the OptiChamber software after any stage of installation:

- Press the button **Start** at the Windows Taskbar, then choose **Settings | Control Panel** from the pop-up menu.
- In the **Control Panel** window open the item **Add/Remove Programs**.
- Click **OptiChamber2** in the program list of the panel **Install/Uninstall** and press the button **Add/Remove**.

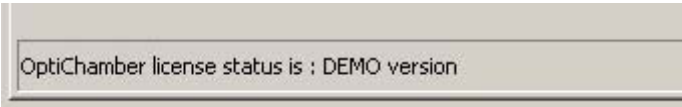
or:

- Select **OptiChamber2 | UnInstall OptiChamber2** at the **Start/Programs** menu.

3. Registration and Installation of the Full-Operation Copy

3.1. Registration

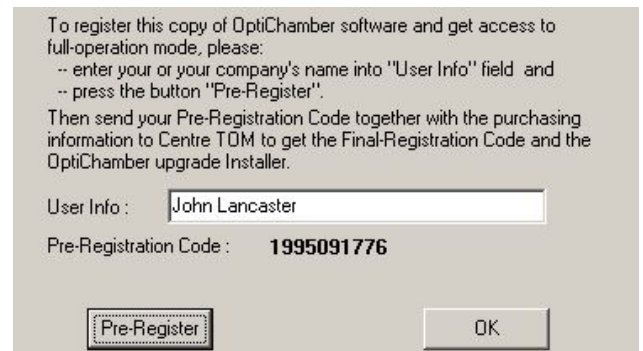
After the first-step installation the user gets a demonstration copy only. This is indicated on the status line at the bottom of the OptiChamber window:



At this stage you are unable to work with real optical chamber parameters that you enter into the data fields except for the number of rays being traced, and the laser rod parameters. All other data have some preset values that remain unchanged regardless of your input.

To turn this demo version into a full-operation package, you first should obtain your personal Pre-Registration Code. Open the item Help/Register of the main menu. This brings about the OptiChamber Registration form. Follow the instructions and enter your personal information into the User Info form (for example, **John Lancaster**). Then click the Pre-Registration button and read the numerical code that appears in the field Pre-Registration Code (e.g., **1995091776**). Send this Pre-Registration Code along with your Purchase Order to DDC Technologies, Inc.

DDC Technologies, Inc.
2980 Waverly Avenue,
Oceanside, NY 11572
Phone: (516) 764-1687
Fax: (516) 764-4412
E-mail: ddctech@yahoo.com



3.2. Final Registration

In response the Developer will email you the payment requisites. After the payment you will receive by email the Final (Second-Step) Installer (**OC2_Register.exe**) and your personal Final-Registration Serial Number (for example, **12345663**), which is needed to launch the setup program. Follow once more the procedure of item 2.2 and run the second-step installation. This time you must enter the Final-Registration Serial Number. On completing the installation you get a full-operation copy of the OptiChamber v.2.0 software.



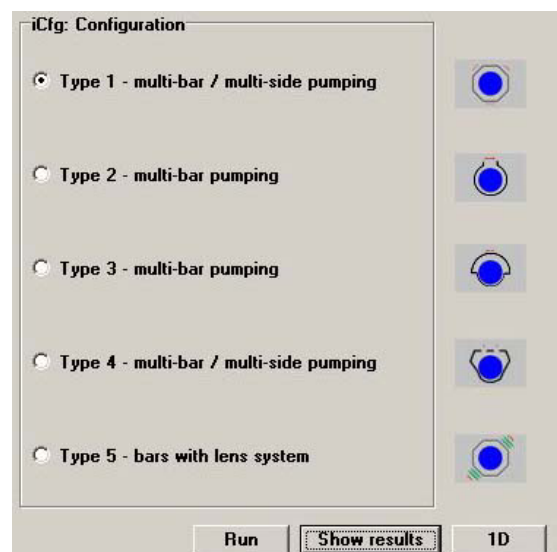
4. Getting started

4.1. Setting desired configuration of the pumping system

Run **OptiChamber.exe** from the corresponding program group of the **Start/Programs** menu or from the directory to which it was installed. Being properly installed, the program starts with the message "**OptiChamber license status is OK**" on the status bar at the bottom of the panel **General**. Choose the desired configuration type from the panel **General**, group **iCfg: Configuration**. After that all the panels which are necessary for definition of the modeling parameters, - **General**, **Diode**, **Rod**, **Cfg 1**, - becomes accessible.

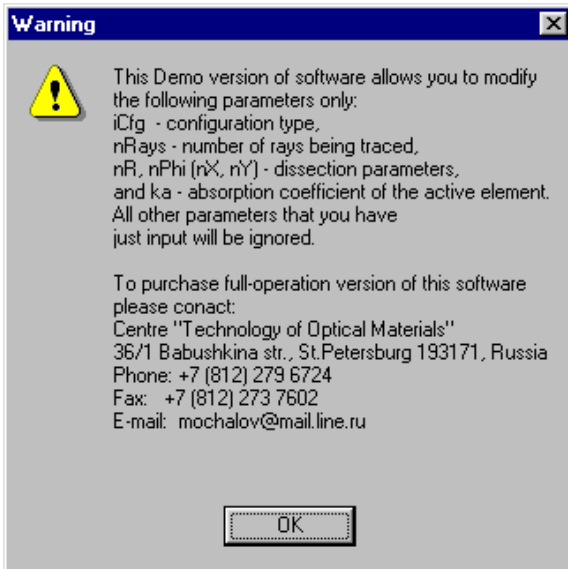
4.2. Starting calculations

To look how the software works, you can start calculation of the pumping system characteristics immediately with predefined parameters by clicking the button **Run** on the panel **General**. This starts the console application, or kernel, **OCCalc2.exe** performing



all necessary calculations, the status bar being set to "Starting...", then to "Calculations in progress, please wait...". To save the screen space, the program runs in minimized-window mode and appears at the Windows taskbar only.

If you are running the Pre-Registration copy instead of the full-operation software, the Demo-version kernel, **OCCalc2Demo.exe**, starts after the warning message.



4.3. Calculation progress

At present, only the number of traced rays that is output in the kernel (**OCCalc2.exe**) window is indication of the kernel progress. OptiChamberCalc.exe window is DOS-styled, with black background. It first appears as an icon (a rectangular with the title **OCCalc2**) at the Windows95/98/2000 taskbar when the Run button is pressed. It may be opened with the mouse by a double-click. The window shows the progress of the *Monte-Carlo* procedure in the form

N = <number of rays being already traced>.

Having total number of the traced rays shown on OptiChamber panel **General**, the user can easily define how much work is done.

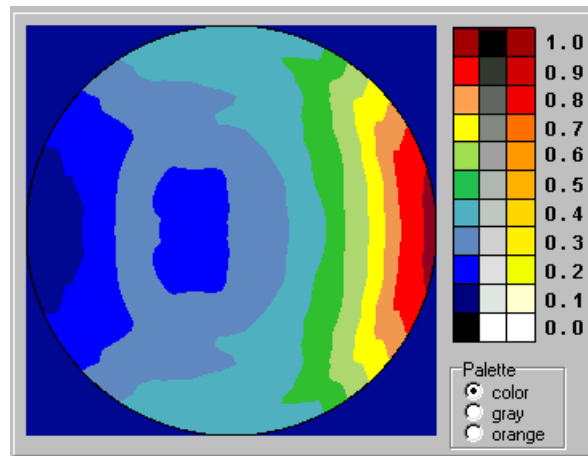
You can **abort calculations** by double-clicking the **OCCalc2** icon at the Windows95/98/2000 taskbar and closing the OptiChamberCalc's window with the help of the X button at the right top of this window.

When the calculations finish, **OCCalc2** quits, the status bar is "Calculations are finished. Press [Show results] to visualize", and two buttons, **Show results** and **1D**, become active.

4.4. Visualizing results

Click on Show results button to build two-dimensional plot of the absorbed power density distribution over area of active medium (the lamp is located at the right side of the rod). Status bar changes to "Drawing...", then, after the plot is created, to the results set containing:

- current configuration type *iCfg*,
- number of radial layers *nR* and angular slices *nPhi* of the active element dissection,



iCfg=2, *nR*=20, *nPhi*=20, *spp*=0.3 W/cm, Absorbed power = 0.1644 W/cm, Efficiency = 54.79 %, *Wmax* = 3.308 W/cm³

- total lineal power of the pumping pulse emitted by the lamp or diode *Plp* [W/cm],
- total lineal power absorbed in the active medium [W/cm],
- efficiency, that is, the ratio of total power absorbed in the active medium to total power emitted by the lamp or diode [percents].
- maximum value of the absorbed power density *Wmax* [W/cm³].

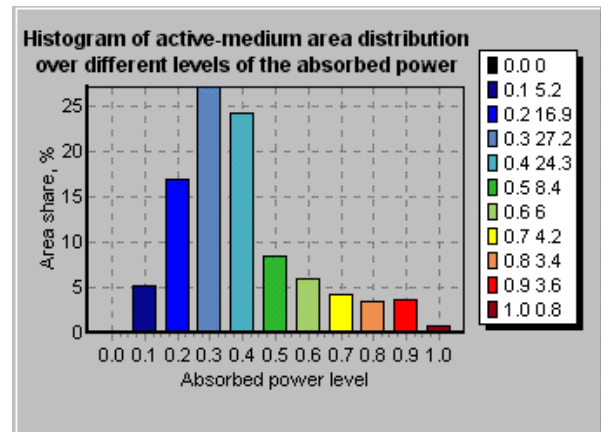
When the mouse pointer enters the plot area, it changes to a cross. Simultaneously, the status bar content changes to show parameters of the point under the mouse pointer:

X = -0.005 cm, Y = 0.003 cm, R = 0.006 cm, Angle = 146.000 degrees, *Wrel* = 0.755, *Wabs* = 2.090 W/cm³

- physical coordinates of the point with respect to the rod center (bottom left corner in case of rectangular rod) *X* and *Y* [cm],
- radial coordinate of the point *R* [cm], $R^2 = X^2 + Y^2$,
- azimuth angle of the point, degrees,
- ratio of the local specific absorbed power density to its maximum value over the rod cross section $Wrel \in [0,1]$,
- absolute value of the local absorbed power density *Wabs* [W/cm³].

Radio group **Palette** provides switching between **Color**, **Gray**, and **Orange** palettes which are used for marking areas with the same range of the absorbed power. There are eleven ranges whose colors are indicated in the scale on the right of the plot, representing the following grades of the relative absorbed power density W_{rel} :

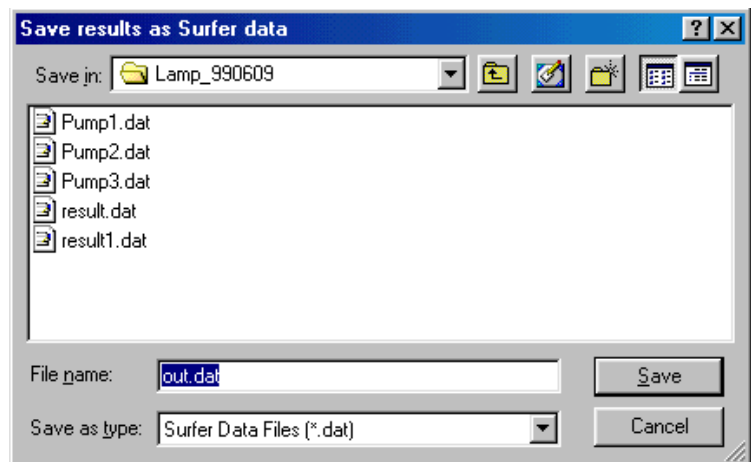
range label	range definition
0.0	$0.00 \leq W_{rel} \leq 0.05$
0.1	$0.05 < W_{rel} \leq 0.15$
0.2	$0.15 < W_{rel} \leq 0.25$
0.3	$0.25 < W_{rel} \leq 0.35$
0.4	$0.35 < W_{rel} \leq 0.45$
0.5	$0.45 < W_{rel} \leq 0.55$
0.6	$0.55 < W_{rel} \leq 0.65$
0.7	$0.65 < W_{rel} \leq 0.75$
0.8	$0.75 < W_{rel} \leq 0.85$
0.9	$0.85 < W_{rel} \leq 0.95$
1.0	$0.95 < W_{rel} \leq 1.00$



The button **1D** allows passing to another representation of the data obtained: a histogram. Being pressed, this button replaces the 2D plot by its one-dimensional counterpart in which the area share of each absorbed power level is plotted versus the value of the relative absorbed power density W_{rel} . Next click on the same button (now its caption is **2D**) brings us back to two-dimensional picture.

4.5. Saving results

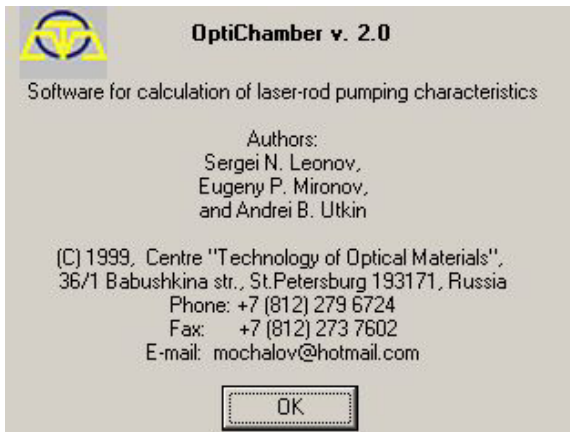
You can save the obtained results on disk with **Results / Save As** item of the main menu. Three submenus - **OptiChamber**, **MathCAD**, and **Surfer data** - offer to save resulting figures and plots in OptiChamber native (*.dat file), MathCAD (*.prn file) and Surfer (*.dat file)



scientific-graphics package formats respectively. The last menu **Bitmap** enables to save 2D plot in bitmap format (*.bmp file). *Warning: bitmap format of the saved pictures has extended features that sometimes cannot be properly recognized by popular graphic editors and viewers (for example, Adobe PhotoShop). If the picture looks distorted in your graphics package, please open the bitmap file with the help of Windows Paint (that is a part of Windows 95/98 package that can be launched from the Start/Programs menu, program group Accessories) and, then, save it back to the disk. After that the bitmap is decoded properly by all packages.* In all cases the standard Windows 95 Save File dialog will guide you through the save procedure, including choice of drive, folder, and file name.

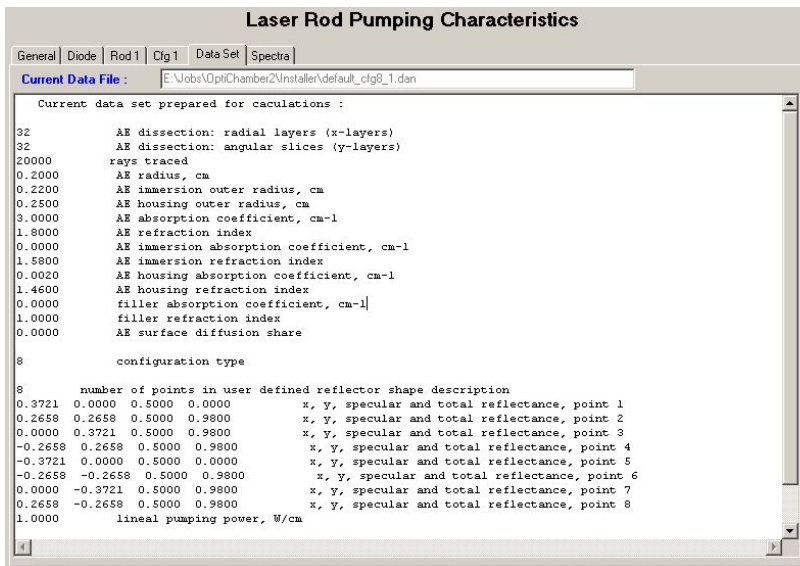
4.6. Loading previously saved results

Results that were previously saved with **Results / Save As / OptiChamber data** option may be read and re-converted into plots with the help of **Results / Load From File** item of the main menu. Corresponding graphics is created just after you close standard Windows Open File dialog window (that enables to choose desired file), so there is no need to click on **Show results** button.



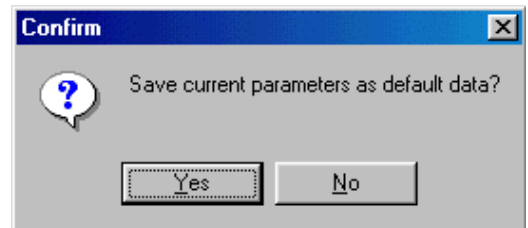
4.7. Getting help

Current version of the software package does not support on-line help. Instead, some comments to the particular program units are added as hints that appear as the mouse pointer enters the unit's area. In addition, this User's Guide can be read directly from the program shell with **Help / OptiChamber Info** item of the main menu (however, illustrations cannot be shown, so you are strongly encouraged to use Microsoft Word instead). Use **Help / Spectral Data Summary** and **Help / About** item to display information about the reference spectra files and the authors.



4.9. Exiting program

To exit to the program, use the standard **File / Exit** item of the main menu. Before quitting, the program prompts you to save data that you entered into the interface fields during current session. If you click on **Yes**, the parameters you defined will be written into the file **pump.ini** and retrieved during the next start-up, otherwise the startup data file remains unchanged.

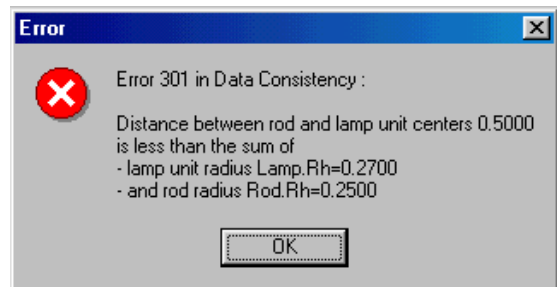


5. Worksheet General: Changing General Parameters

5.1. Input fields

Now, when you have learned how to start calculations and exit the program, let us explain how to change parameters of the optical system under modeling. This should be done by changing values in corresponding *input fields*. Each input field is supplied by *up-down control* consisting of a pair of arrow buttons, allowing users to change the size of a numerical value by

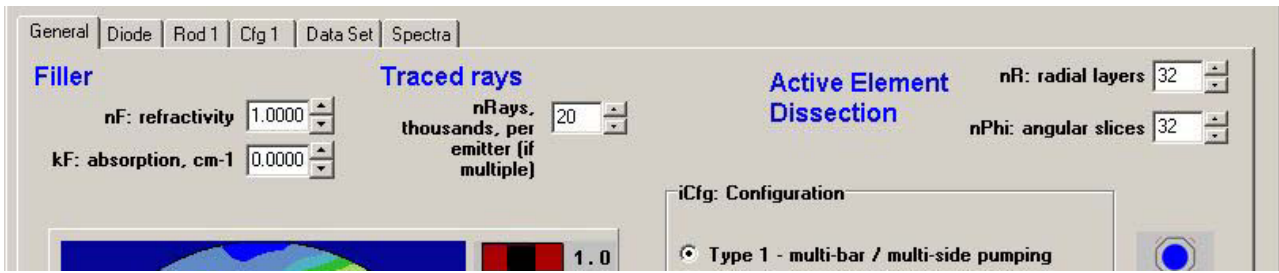
clicking on them. The step of increase and decrease is predefined for each field and the excursion length is confined in accordance with the physical meaning of the parameter in question and current values of other parameters. For example, it is possible to decrease the refraction indices down to the value 1 only and it is impossible to make, say, the rod radius greater than the current radius of surrounding reflector. If you prefer to be free of finite-step and data-consistency restrictions, you can enter in the input field any desired value using a keyboard. However, when you start calculations (see section 4.2) the program shell first performs the data-consistency check and in case of any data mismatch opens pop-up window with error diagnostics rather than starts the kernel file **OCCalc2.exe**. All input fields are subdivided into groups representing closely related parameters: General, Lamp/Diode, Laser Rod, and Configuration. Groups are placed in separate worksheets, each of which can be brought to front by clicking on **General**, **Diode**, **Laser Rod**, or **Config n** bookmark at the top row.



On the worksheet **General**, apart from the **Configuration** radio group allowing to choose desired pumping system configuration type *iCfg*, there are five input fields in three local subgroups. The subgroup **Filler** provides opportunity to insert two optical characteristics of the medium in which the entire pumping system is embedded: the **absorption coefficient** kF [cm^{-1}] and the **refraction index** nF . As in most cases this medium is air, the default values of these parameters are zero and unity respectively. The subgroup **Traced rays** consists of a single field in which one can change number of rays $nRays$ being generated by the kernel in the course of the *Monte-Carlo* procedure. The subgroup **Active Element Dissection** includes two input fields for the number of radial layers nR and number angular slices $nPhi$ to subdivision of the active-medium area into finite sample elements by the polar-coordinate dissection.

4.8. Getting auxiliary information

To get information about the parameters that are currently set as the initial data for the *kernel* program **OCCalc2.exe**, bring to front the **Data Set** worksheet clicking on corresponding bookmark at the top bookmark row. The upper field **Current Data File** contains the name of resent file (**pump.ini** by default) from which the initial data were loaded (see section 4.5). Below one can read full list of currently active parameters prepared for the next-in-turn calculation run. In the next sections we explain how to change each of these parameters with the help of interface fields on other worksheets.



6. Worksheet Diode

Use this worksheet to modify parameters of the pumping source.

6.1. Worksheet Diode subgroups

There are two subgroups in this worksheet: **Lineal Pumping Power**, **Beam Divergence** and **Spectral Properties**. **Lineal Pumping Power** let you enter the power of the pumping pulse that is emitted by unit-length segment of the diode bar. The table **Spectral Properties** allows user to enter spectral characteristics of the emitting diode.

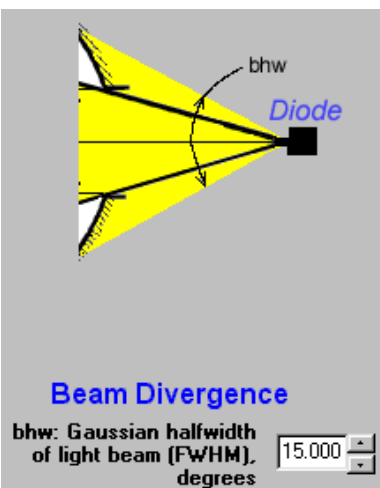
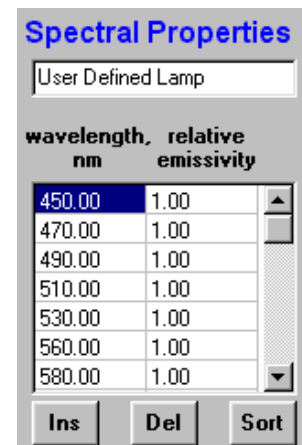
6.2. Worksheet Diode, subgroup Lineal Pumping Power

Here you can enter the power P_{lp} of the pumping source. As far as the two-dimensional approximation is used that implies the infinitely long diode bar, the power is measured in Watts per unit (1 cm) of the diode bar length (so-called lineal power). Since all energy relations are linear, you can treat P_{lp} as the total diode bar power (implying uniform distribution of radiation along the lamp length) or, furthermore, as the total diode bar energy (implying constant value of the power for all duration of the pumping pulse).



6.3. Worksheet Diode, subgroup Spectral Properties

The software in its present form uses the one-spectral-band approximation that actually means that all rays are generated in the *Monte-Carlo* procedure at the same wavelength. Thus, the model fails to reproduce spectral variations of the pumping system parameters. The only thing that can be done in the framework of the one-band approximation is to calculate a rough estimation for the effective absorption coefficient of the laser rod ka for given combination of the lamp emission and rod absorption spectra. Initial data for this procedure (to be discussed in section 7.2) concerning the diode spectral properties should be entered into the *Spectral Properties* table. The table consists of the editable title field (containing user-defined reference information) and two columns: wavelength [nm] and diode emissivity. The final absorption coefficient is normalized on the integral diode emissivity, so you can enter the diode emissivity in any desired units. You can insert and delete single lines of the table using the **Ins** and **Del** buttons beneath. To sort the table in the ascending-wavelength order, use the **Sort** button. Input data (together with the reference information in the title field) can be loaded from or saved to a file with **Load Diode Spectrum** and **Save Diode Spectrum** items of the section **Diode** of main menu. Default extension for the diode-spectrum data file is *.ls. Several files containing standard diode spectra are included with corresponding comments in the present package as a reference information. Use **Help / Spectral Data Summary** item of the main menu to see the list of all diode-spectrum files and their brief descriptions.



6.4. Worksheet Diode, subgroup Beam Divergence

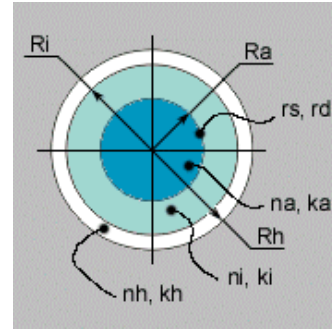
A diode with the Gaussian beam distribution is assumed as a pumping source. Full width of this distribution at half maximum, bhw [degrees], is defined by user in **Beam Divergence** subgroup.

7. Worksheet Rod

This worksheet allows to modify parameters of the laser rod to be pumped.

7.1. Worksheet Rod subgroups

All configurations include cylindric laser rods. Optional immersion and housing (outer protection) layers are included. Worksheet subgroups are: **Active Element**, **Active Element Surface**, **Immersion**, **Housing**, and **Spectral Properties**. Here other parameters appear is the **Active Element** subgroup while **Immersion** and **Housing** subgroups do not present.



7.2. Worksheet Rod, subgroup Active Element

The subgroup contains parameters of the active-medium area:

- radius Ra [cm],
- refraction index na ,
- absorption coefficient ka [cm^{-1}].

In the case of configuration 4 the first parameter is replaced by two - wa [cm] and ha [cm] - characterizing size of the rectangular rod.

The button **Calculate from spectra** allows calculating effective absorption coefficient ka on the basis of diode and rod spectra that is input in the **Spectral Properties** tables of the **Diode** and **Rod** worksheets. First the program defines the spectral area Λ which is the intersection lamp and rod spectra. Then the effective absorption coefficient ka is calculated according to the expression

$$ka = ddf \int_{\Lambda} \sigma(\lambda) k(\lambda) d\lambda / \int_{\Lambda} \sigma(\lambda) d\lambda$$

where ddf is a dopant density factor that you can enter into the field below ka input field, $\sigma(\lambda)$ is the diode emissivity measured in some conventional units and $k(\lambda)$ is the absorption coefficient for the wavelength λ . The default value of the dopant density factor ddf is unity, so it does not affect the resulting value of ka . However, if you have to calculate the absorption coefficient for the dopant density that differ from that of the spectral table, you should calculate ddf as a ratio of the actual dopant density to that for which the spectrum table is calculated. In this case the correct value of the absorption coefficient is calculated without need of re-insertion of the scaled absorption coefficients throughout the entire spectrum.

Finally, the obtained value is inserted into the absorption coefficient field. In integrating, linear data interpolation is used, so in case of linear spectrum one should use zero limiting values to mark separate spectral lines. That is, the set

```
450 200
670 200
449 0
450 200
451 0
669 0
670 200
671 0
```

corresponds to two narrow spectral lines.

7.3. Worksheet Rod, subgroup Active Element Surface

Optical properties of the active element surface, which can be coarse due to special treatment, are characterized by two parameters:

- probability of ray refraction/reflection rs and
- probability of ray isotropic random scattering rd .

7.4. Worksheet Rod, subgroup Immersion

This subgroup is the same as that of worksheet Lamp (see section 6.3) and contains the following parameters:

- outer radius Ri [cm],
- refraction index ni ,
- absorption coefficient ki [cm^{-1}].

Immersion layer can be excluded by setting zero thickness ($Ri = Ra$).

Active Element

Ra: radius, cm

na: refractivity

ka: absorption, cm-1

Calculate from spectra

ddf: dopant density factor

Active Element Surface

rs: probability of ray refraction/reflection

rd: probability of ray random scattering

Immersion

Ri: outer radius, cm

ni: refractivity

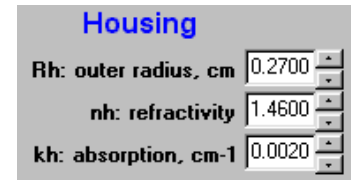
ki: absorption, cm-1

7.5. Worksheet Rod, subgroup Housing

The housing parameters are:

- outer radius Rh [cm],
- refraction index nh ,
- absorption coefficient kh [cm^{-1}].

Housing layer can be excluded by setting zero thickness ($Rh = Ri$).

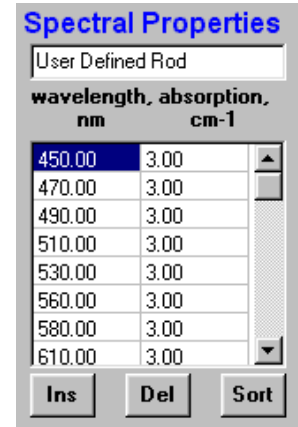


The screenshot shows a dialog box titled "Housing" with three input fields, each with a spin button on the right:

- Rh: outer radius, cm: 0.2700
- nh: refractivity: 1.4600
- kh: absorption, cm-1: 0.0020

7.6. Worksheet Rod, subgroup Spectral Properties

The table **Spectral Properties** enables the user to enter dependence of the absorption coefficient of the active medium on the wavelength, which is used by **Calculate from spectra** procedure (see sections 6.6 and 7.2). It consists of the editable title field (containing user-defined reference information) and two columns: wavelength [nm] and absorption [cm^{-1}]. You can insert and delete single lines of the table using the **Ins** and **Del** buttons. To sort the table in the ascending-wavelength order, use the **Sort** button. Input data (together with user-defined reference information of the title field) can be loaded from or saved to a file with **Load Rod Spectrum** and **Save Rod Spectrum** items of the section **Rod** of main menu. Default extension for the rod-spectrum data file is *.rs. Several files containing standard rod absorption spectra are included with corresponding comments in the present package as a reference information. Use **Help / Spectral Data Summary** item of the main menu to see the list of all rod-spectrum files and their brief descriptions.



The screenshot shows a dialog box titled "Spectral Properties" with a title field and a table:

User Defined Rod

wavelength, nm	absorption, cm^{-1}
450.00	3.00
470.00	3.00
490.00	3.00
510.00	3.00
530.00	3.00
560.00	3.00
580.00	3.00
610.00	3.00

Buttons: Ins, Del, Sort

8. Configuration Worksheets

Parameters that are related to diode bars and rod position and the reflector dimensions are placed on five worksheets named as **Cfg 1**, one worksheet per each pre-defined pumping-system configuration type. Only one worksheet is active at a time - one that is chosen in *iCfg* radio group on the worksheet **General**.

The **reflective surfaces** are characterized by two parameters: *rd* (**diffuse reflectance**) and *rt* (**total reflectance**). Sample rays hitting the surface are subjected to: uniform back scattering within the solid angle 2π with probability *rd* (diffuse reflectance), specular reflection with probability $rs = rt - rd$, and absorption with probability $1-rt$.

It should be noted that the number of diode bar groups both with diode parameters and also reflector shape peculiarities may be subject of change for each of the pre-defined configurations.

System composition and subgroups for each configuration are listed below. Configuration parameters are listed in Appendix A.

8.1. Pre-defined configuration 1

Composition: laser rod, diode bars, and hexagonal reflector with input windows.

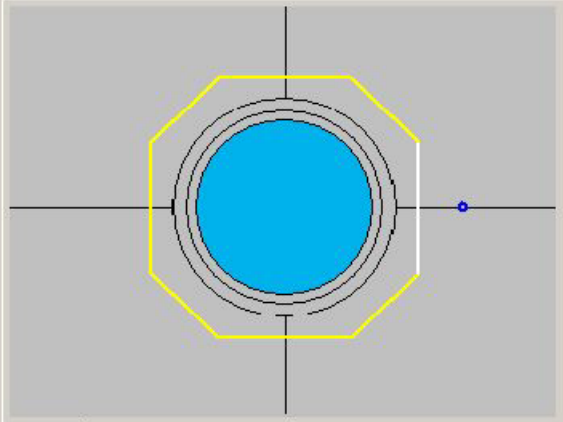
Subgroups: Diode bar parameters controls, Reflector Shape, Reflectances, Diode Bar Groups, Scale Shape, Move Bars and layout sketch Panels.

Coordinates of the reflector polygon vertexes and reflectances of corresponding segments are defined by user in the table at the right side of panel both with diode bar groups location. When you are editing the table, the changes are reflected in the sketch at the left, current point or segment being marked by color. The origin of coordinates coincides with the laser rod center. You can insert and delete single lines of the table using the **Ins** and **Del** buttons beneath. Input data can be loaded from or saved to a file with **Load** (in this case the filename appears below the **Configuration** radio group on the worksheet **General**) and **Save** buttons. Default extension for the diode-spectrum data file is *.shp. Using Scale Shape and Move Bars panels you can control extensions and position of whole reflector shape and also diode bars location.

Laser Rod Pumping Characteristics

General | Diode | Rod 1 | **Cfg 1** | Data Set | Spectra

Number of diode bar groups:
 Number of diode bars in each group:
 Emitting length for each diode, cm:
 Diode bars step, cm:



Reflector Shape		Reflectances		Diode Bar Groups		
x, cm	y, cm	diffuse	total	bar exist	x_centre, cm	y_centre, cm
0.1500	0.3000	0.0000	0.9800	0		
-0.1500	0.3000	0.0000	0.9800	0		
-0.3000	0.1500	0.5000	0.9800	0		
-0.3000	-0.1500	0.0000	0.9800	0		
-0.1500	-0.3000	0.0000	0.9800	0		
0.1500	-0.3000	0.0000	0.9800	0		
0.3000	-0.1500	0.0000	0.0	1	0.400	0.0
0.3000	0.1500	0.0000	0.9800	0		

Row Table

Scale Shape: / x

Move Bars X

Reflector Shape (total refl. > 50%)
 Reflector Shape (total refl. <= 50%)
 Diode Bars
 Laser Rod

8.2. Pre-defined configuration 2

Composition: laser rod, diode bar, and cylindrical reflector with flat window.

Subgroups: Diode bar parameters controls, Reflector Shape, Reflectances, Diode Bar Groups, Scale Shape, Move Bars and layout sketch Panels.

Laser Rod Pumping Characteristics

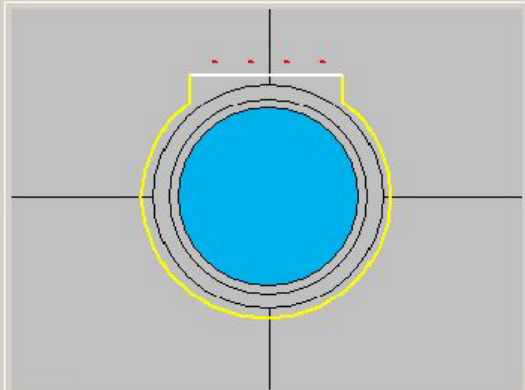
General | Diode | Rod 1 | Cfg 1 | Data Set | Spectra

Number of diode bar groups:

Number of diode bars in each group:

Emitting length for each diode, cm:

Diode bars step, cm:



█ Reflector Shape (total refl. > 50%)
█ Reflector Shape (total refl. <= 50%)
█ Diode Bars
█ Laser Rod

Reflector Shape		Reflectances		Diode Bar Groups		
x, cm	y, cm	diffuse	total	bar exist	x_centre, cm	y_centre, cm
0.2612	0.0609	0.0000	0.9800	0		
0.2410	0.1187	0.0000	0.9800	0		
0.2085	0.1701	0.0000	0.9800	0		
0.1652	0.2132	0.0000	0.9800	0		
0.1652	0.2730	0.0000	0.0100	1	0.000	0.302
-0.1752	0.2730	0.0000	0.9800	0		
-0.1752	0.2132	0.0000	0.9800	0		
-0.2185	0.1701	0.0000	0.9800	0		
-0.2510	0.1187	0.0000	0.9800	0		

Scale Shape: / x

Up Right

Down

Ins Del Load Save
Row Table

Move Bars X Closer Remotely

8.3. Pre-defined configuration 3

Composition: laser rod, diode bar, and reflector consisting of two semi-cylinders adjoined by flat surface.

Subgroups: Diode bar parameters controls, Reflector Shape, Reflectances, Diode Bar Groups, Scale Shape, Move Bars and layout sketch Panels.

Laser Rod Pumping Characteristics

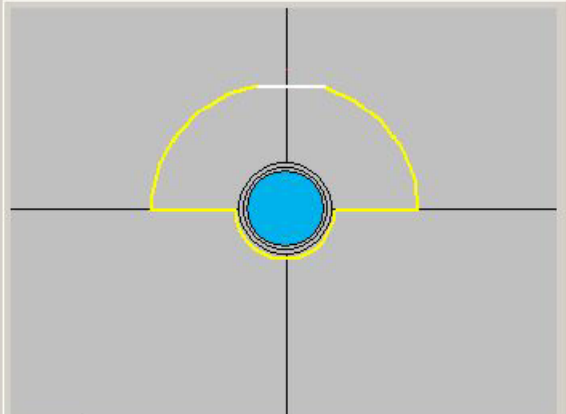
General | Diode | Rod1 | Cfg1 | Data Set | Spectra

Number of diode bar groups:

Number of diode bars in each group:

Emitting length for each diode, cm:

Diode bars step, cm:



Reflector Shape		Reflectances		Diode Bar Groups		
x, cm	y, cm	diffuse	total	bar exist	x_centre, cm	y_centre, cm
0.6950	-0.0100	0.5500	0.9800	0		
0.6730	0.1641	0.5500	0.9800	0		
0.6084	0.3272	0.5500	0.9800	0		
0.5053	0.4692	0.5500	0.9800	0		
0.3701	0.5810	0.5500	0.9800	0		
0.2113	0.6557	0.5500	0.0000	1	0.000	0.750
-0.1538	0.6676	0.5500	0.9800	0		
-0.3031	0.6234	0.5500	0.9800	0		
-0.4512	0.5294	0.5500	0.9800	0		

Reflector Shape (total refl. > 50%)
 Reflector Shape (total refl. <= 50%)
 Diode Bars
 Laser Rod

Scale Shape: / x

Up Down

Left Right

Ins Del Load Save
Row Table

Move Bars X Remotely

Closer

8.4. Pre-defined configuration 4

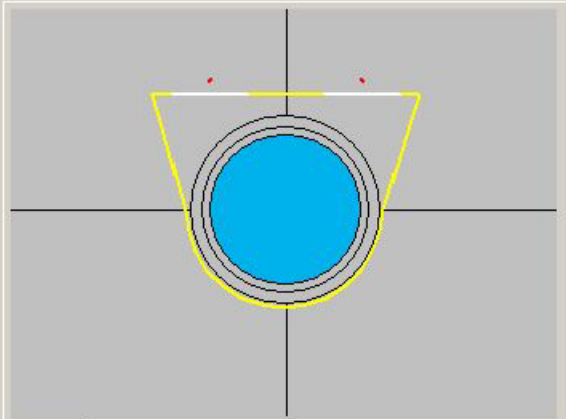
Composition: laser rod, two diode bars, quasi-parallelepiped reflector confined at the lamp side by adjoined semi-cylinder, and filter plate.

Subgroups: Diode bar parameters controls, Reflector Shape, Reflectances, Diode Bar Groups, Scale Shape, Move Bars and layout sketch Panels.

Laser Rod Pumping Characteristics

General | Diode | Rod 1 | Cfg 1 | Data Set | Spectra

Number of diode bar groups:
 Number of diode bars in each group:
 Emitting length for each diode, cm:
 Diode bars step, cm:



Reflector Shape		Reflectances		Diode Bar Groups		
x, cm	y, cm	diffuse	total	bar exist	x_centre, cm	y_centre, cm
0.3500	0.3100	0.5500	0.9800	0		
0.3000	0.3100	0.0000	0.0000	1	0.200	0.350
0.1000	0.3100	0.5500	0.9800	0		
-0.1000	0.3100	0.0000	0.0000	1	-0.200	0.350
-0.3000	0.3100	0.5500	0.9800	0		
-0.3500	0.3100	0.5500	0.9800	0		
-0.2650	0.0000	0.5500	0.9800	0		
-0.2585	-0.0580	0.5500	0.9800	0		
-0.2393	-0.1130	0.5500	0.9800	0		

Scale Shape: / 1.10 x

Up: 0.010 Down

Left Right

Ins Del Load Save
Row Table

Move Bars X 1.01 Remotely

Reflector Shape (total refl. > 50%)
 Reflector Shape (total refl. <= 50%)
 Diode Bars
 Laser Rod

8.5. Pre-defined configuration 5

Composition: laser rod, diode bars, hexagonal reflector with input windows, and lens systems.

Subgroups: Diode bar parameters controls, Reflector Shape, Reflectances, Diode Bar Groups, Scale Shape, Move Bars, lens system, and layout sketch panels.

Laser Rod Pumping Characteristics

General
Diode
Rod1
Cfg1
Data Set
Spectra

Number of diode bar groups

Number of diode bars in each group

Emitting length for each diode, cm

Diode bars step, cm

First lens thickness, cm

Gap between lenses, cm

Second lens thickness, cm

Dist. rod center/first lens, cm

Lenses diameter, cm

Reflector-lens gap :

refraction index

absorption coefficient, cm-1

First lens :

R1, cm

R2, cm

abs. index, cm-1

refr. index

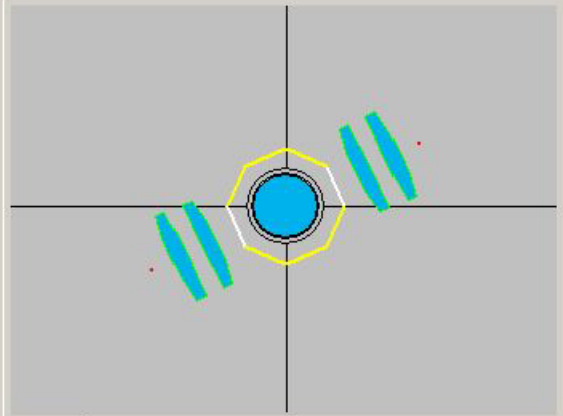
Second lens :

R1, cm

R2, cm

abs. index, cm-1

refr. index



Reflector Shape		Reflectances		Diode Bar Groups		
x, cm	y, cm	diffuse	total	bar exist	x_centre, cm	y_centre, cm
0.3721	0.0000	0.5000	0.0000	1	0.846	0.412
0.2658	0.2658	0.5000	0.9800	0		
0.0000	0.3721	0.5000	0.9800	0		
-0.2658	0.2658	0.5000	0.9800	0		
-0.3721	0.0000	0.5000	0.0000	1	-0.846	-0.412
-0.2658	-0.2658	0.5000	0.9800	0		
0.0000	-0.3721	0.5000	0.9800	0		
0.2658	-0.2658	0.5000	0.9800	0		

Scale Shape

/ x

Up

Left Right

Down

Row Table

Move Bars X

Reflector Shape (total refl. > 50%)

Reflector Shape (total refl. <= 50%)

Diode Bars

Laser Rod

9. Main Menu

Pull-down main menu is located at the top of the OptiChamber window. It consists of five top-level items: **File**, **Lamp**, **Rod**, **Results**, and **Help**. The last four are already described above: **Lamp** in section 6.6, **Rod** in 7.6, **Results** in 4.4 and 4.5, and **Help** in 4.6. The item **File** consists of five sub-items:

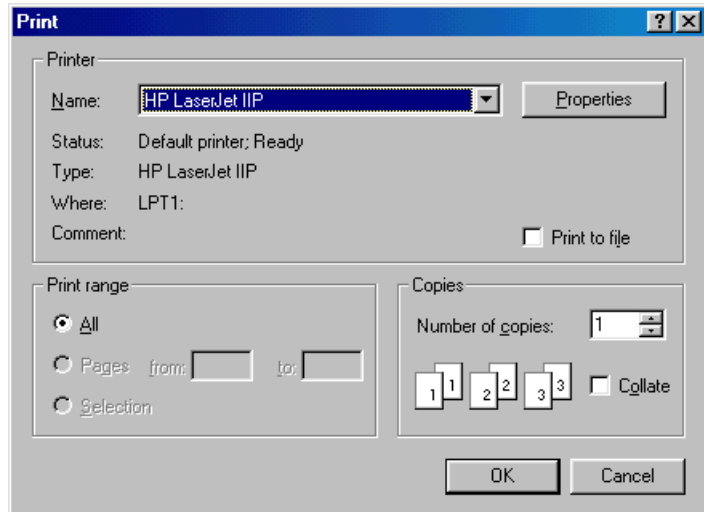
- **Open Chamber** allows you to open previously saved data that were input in **OptiChamber** interface shell. In this case corresponding file name will appear on the right in the window title.



- **Save Chamber** lets you to save the input data on file (default file extension is *.dan). It differs from **Save As** item of the **Results** menu in that the latter saves *results* of work of the kernel program **OCCalc2** while here the *initial parameters* are saved.

- **Save Chamber as Default** enables to write the input data into **pump.ini** file. The parameters you entered will be retrieved at the next start-up.

- **Printer setup** opens standard Windows Print Setup Dialog in which the user can setup desired printer and its parameters.
- **Print results** allows to print the plot of two-dimensional absorbed power density distribution (see section 3.3) together with the list of initial parameters. It brings about standard Windows Print Dialog in which you can setup desired printer and its options.



- **Exit** closes the program OptiChamber. Before quitting, the program prompts you to save data that you entered into the interface fields during current session (see section 3.8). If you click on **Yes**, the parameters you defined will be written into the file **pump.ini** and retrieved during the next start-up, otherwise the startup data file remains unchanged.

Appendix A. List of Parameters

G. General

G.1. Filler

G.1.1. Filler absorption coefficient (kF), [cm⁻¹]

G.1.2. Filler refractivity (nF)

G.2. Traced rays

G.2.1. Number of rays being traced (nRays)

G.3. Active Element Dissection

G.3.1.1. Radial layers (nR)

G.3.2.1. Angular slices (nPhi)

G.4. Configuration type

G.4.1. Number indicating configuration type (iCfg) [1..5]

L. Lamp (all configurations except 6)

L.1. Envelope

L.1.1. Inner radius (Rl), [cm]

L.1.2. Outer radius (Re), [cm]

L.1.3. Refractivity (ne)

L.1.4. Absorption (ke), [cm⁻¹]

L.2. Immersion

L.2.1. Outer radius (Ri), [cm]

L.2.2. Refractivity (ni)
L.2.3. Absorption (ki), [cm-1]

L.3. Housing

L.3.1. Outer radius (Rh), [cm-1]
L.3.2. Refractivity (nh)
L.3.3. Absorption (kh), [cm-1]

D. Diode

D.1. Beam divergence

D.1.1. Gaussian halfwidth of light beam, FWHM, (bhw), [degrees]

D.2. Spectral properties

D.2.1. Wavelength [nm]
D.2.2. Relative emissivity

D.3. Lineal pumping power

D.3.1. Lineal pumping power, (plp), [W/cm]

R. Laser Rod

R.1. Active Element

R.1.1. Radius (Ra), [cm]
R.1.2. Refractivity (na)
R.1.3. Absorption (ka), [cm-1]
R.1.4. Dopant density factor (ddf)

R.2. Ray Interaction with Active Element Surface

R.2.1. Probability of ray refraction/reflection (rs)
R.2.2. Probability of ray random scattering (rd)

R.3. Immersion

R.3.1. Outer radius (Ri), [cm]
R.3.2. Refractivity (ni)
R.3.3. Absorption (ki), [cm-1]

R.4. Housing

R.4.1. Outer radius (Rh), [cm]
R.4.2. Refractivity (nh)
R.4.3. Absorption (kh), [cm-1]

R.5. Spectral Properties

R.5.1. Wavelength [nm]
R.5.2. Absorption [cm-1]

1. Configurations 1 to 4

1.1. Diode bars parameters

1.1.1. Number of diode bar groups
1.1.2. Number of diode bars in each group
1.1.3. Emitting length for each diode, [cm]
1.1.4. Diode bars step, [cm]

1.2. Reflector shape, reflectances, and diode bar groups

1.2.1. X coordinate of the shape segment origin (x), [cm]
1.2.2. Y coordinate of the shape segment origin (y), [cm]
1.2.3. Specular reflectance of the segment (rs)
1.2.4. Total reflectance of the segment (rt)
1.2.5. Presence of diode bar adjacent to current segment, 1 - if the adjacent diode bar exist for this segment, empty - otherwise
1.2.6. X-location of the centre of diode bar group (x_center), [cm]
1.2.7. Y-location of the centre of diode bar group (y_center), [cm]

- 1.3. Reflector shape and bars controls
 - 1.3.1. Multiplier to scale reflector shape uniformly up or down
 - 1.3.2. Shift value [cm] to move entire reflector shape up/down/left/right
 - 1.3.3. Multiplier to scale the position of the central point of diode bar group closer/remotely relative the origin

5. Configuration 5

- 5.1. Diode bars parameters
 - 5.1.1. Number of diode bar groups
 - 5.1.2. Number of diode bars in each group
 - 5.1.3. Emitting length for each diode, [cm]
 - 5.1.4. Diode bars step, [cm]

Remarks:

It is assumed that all diodes in a bar group emitting along direction from the diode bar group center to the origin.

- 5.2. Reflector shape, reflectances, and diode bar groups
 - 5.2.1. X coordinate of the shape segment origin (x), [cm]
 - 5.2.2. Y coordinate of the shape segment origin (y), [cm]
 - 5.2.3. Specular reflectance of the segment (rs)
 - 5.2.4. Total reflectance of the segment (rt)
 - 5.2.5. Presence of diode bar adjacent to current segment, 1 - if the adjacent diode bar exist for this segment, empty - otherwise
 - 5.2.6. X-location of the centre of diode bar group (x_center), [cm]
 - 5.2.7. Y-location of the centre of diode bar group (y_center), [cm]

- 5.3. Reflector shape and bars controls
 - 5.3.1. Multiplier to scale reflector shape uniformly up or down
 - 5.3.2. Shift value [cm] to move entire reflector shape up/down/left/right
 - 5.3.3. Multiplier to scale the position of the central point of diode bar group closer/remotely relative the origin

- 5.4. Lens system parameters
 - 5.4.1. System parameters*
 - 5.4.1.1. First (adjacent to the reflector) lens thickness, [cm]
 - 5.4.1.2. Second (placed remotely from the reflector) lens thickness, [cm]
 - 5.4.1.3. Gap length between lenses, [cm]
 - 5.4.1.4. Distance from the laser rod center (the origin) to the first lens, [cm]
 - 5.4.1.5. Lens diameter (this value is applicable to both lenses), [cm]
 - 5.4.1.6. Refraction index for the medium filling the gap between reflector and the first lens
 - 5.4.1.7. Absorption coefficient for the medium filling the gap between reflector and the first lens, [cm⁻¹]
 - 5.4.2. Lens parameters
 - 5.4.2.1. First lens first surface (nearest to the origin) radius**, [cm⁻¹]
 - 5.4.2.2. First lens second surface (located remotely from the origin) radius**, [cm⁻¹]
 - 5.4.2.3. First lens refraction index
 - 5.4.2.4. First lens absorption coefficient, [cm⁻¹]
 - 5.4.2.5. Second lens first surface (nearest to the origin) radius**, [cm⁻¹]
 - 5.4.2.6. Second lens second surface (located remotely from the origin) radius**, [cm⁻¹]
 - 5.4.2.7. Second lens refraction index
 - 5.4.2.8. Second lens absorption coefficient, [cm⁻¹]

Remarks:

*) Lens system assumed to be centered and aligned along the direction from origin to the corresponding diode bar group center.

***) Radius values are positive for the concave lens surfaces, negative otherwise.