sDNA_GH

sDNA is a world leading tool for Spatial Design Network Analysis. sDNA_GH is a plug-in for Grasshopper providing components that run the tools from a local <u>sDNA</u> installation, on Rhino and Grasshopper geometry and data.

sDNA

sDNA is able to calculate Betweenness, Closeness, Angular distance, and many other quantities including custom hybrid metrics, and is able to perform many other advanced functions as well. Please note, for results of a network analysis to be meaningful, it must be ensured that the network is first properly <u>prepared</u>.

sDNA_GH functionality

sDNA_GH:

- Reads a network's polyline Geometry from Rhino or Grasshopper, and Data from any User Text on it.
- Writes the network polylines (formed by one or more polylines) and user Data to a Shapefile.
- Initiates an sDNA tool that processes that shapefile, and e.g. carries out a network preparation or an analysis.
- Reads the shapefile produced by the sDNA tool.
- Displays the results from sDNA by colouring a new layer of new polylines or the original ones.

User manual.

System Requirements.

Software

- 1. Windows 10 or 8.1 (not tested in Windows 11)
- 2. Python 2.7. Please note Iron Python 2.7 does not run sDNA correctly (as incorrect shapefiles are produced).
- 3. sDNA.
- 4. Rhino and Grasshopper (tested in Rhino 7)

Hardware

- 1. 64-bit Intel or AMD processor (Not ARM)
- 2. No more than 63 CPU Cores.
- 3. 8 GB memory (RAM) or more is recommended.
- 4. 1.2 GB disk space.

Installation.

- 1. Ensure you have an installation of <u>Rhino 3D</u> including Grasshopper (versions 6 and 7 are supported).
- 2. Ensure you have an installation of <u>Python 2.7</u> [^0] that can run sDNA correctly from the command line. sDNA_GH runs sDNA from the command line. Command line use of sDNA has been tested with Python versions 2.6 and 2.7. Do not run sDNA with Iron Python 2.7, as invalid shape files may be produced (it is not possible to access the Iron Python shipped with Rhino from the command line, in any case).
- 3. To use sDNA with sDNA_GH, ensure you have an installation of <u>sDNA</u>. sDNA itself may require the 64 bit (x64) Visual Studio 2008 redistributable, available <u>here</u> or <u>here</u>). The closed source "free as in beer" version of <u>sDNA</u> and sDNA+ may also require the 32 bit (x86) Visual Studio 2008 redistributable available or <u>here</u> or <u>here</u> in order to unlock sDNA with a serial number.
- 4. Download sDNA GH.zip from food4Rhino or the sDNA GH releases page on Github.
- 5. Ensure sDNA_GH.zip is unblocked: Open File Explorer and go to your Downloads folder (or whichever folder you saved it in). Right click it and select Properties from the bottom of the menu. Then click on the *Unblock* check box at the bottom (right of *Security*), then click OK or Apply. The check box and *Security* section should disappear. This should unblock all the files in the zip archive. If any files still need to be unblocked, a PowerShell script is provided in the zip file: \sDNA_GH\dev_tools \batch_files\unblock_all_files_powershell.bat[^2] Please do not automatically trust and unblock all software downloaded from anywhere on the internet [^1].
- 6. Open Rhino and Grasshopper.
- 7. In Grasshopper's pull down menus (above the tabs ribbon at the top) click File -> Special folders -> User Objects Folder. The default in Rhino 7 is %appdata%\Grasshopper\UserObjects. Note, this is not the Components Folder used by many other plug-ins (i.e. not %appdata%\Grasshopper\Libraries).
- 8. Copy sDNA_GH.zip to this folder (e.g. it should be at %appdata%\Grasshopper

\UserObjects\sDNA_GH.zip).

- 9. Unzip sDNA_GH.zip to this location (in Windows 10 right click sDNA_GH.zip and select Extract All ..., then click Extract to use the suggested location). In the User Objects folder, a single subfolder called sDNA_GH should have been created.
- 10. Restart Rhino and Grasshopper.
- 11. The sDNA_GH plug in components should now be available under a new "sDNA" tab in the ribbon amongst the tabs for any other plug-ins installed (right of Mesh, Intersect, Transform and Display etc).
- 12. To use sDNA with sDNA_GH, if no preferences are specified, sDNA_GH will search for sDNA and Python 2.7 installations automatically, using the first one of each it finds. To ensure sDNA_GH uses a particular version of sDNA and the correct Python 2.7 interpreter it is recommended on first usage to: -place a Config component on the canvas (the component with a gear/cog icon in Extra). -Specify the file path of the sDNA folder (containing sDNAUISpec.py and runsdnacommand.py) of the sDNA installation you wish to use in the sDNA_folders input. -Specify the file path of the Python 2.7 interpreter's main executable in the python input. -Specify any other options you wish to save and reuse on all projects, if necessary by adding custom input Params with the option's name. -Connect a true boolean toggle to go. An installation wide user options file (config.toml) will be created if there isn't one already. -To save options to other project specific config.toml files, specify the file path in save_to and repeat the previous 4 sub steps.
- 13. For a first test of sDNA_GH using sDNA, open \sDNA_GH\tests \5x18_random_grid_network.3dm (in the previously unzipped folder in the User Objects folder), place an sDNA_Integral component and connect a true boolean toggle to its go.
- 14. If a newer version of sDNA is used in future with tools unknown to sDNA_GH at the time it was built, if a Config component is placed, and the path of the new sDNA specified in sDNA_folders, sDNA_GH will attempt to automatically build components and user objects for the new sDNA tools, and add them to Grasshopper for you. Set make_new_comps to false to prevent this.

Usage.

Components.

Automatic multi-tools.

Each sDNA tool has its own Grasshopper component. To run a tool, a true value, e.g. from a

Boolean toggle component, must be connected to its component's go Input Param [^note]. To group together common work flows, if an auto_ option is set to true, some tools also automatically run other tools before or after they run themselves. For example, this allows an entire sDNA process to be run on Rhino Geometry from a single sDNA tool component. When an sDNA_GH component is first placed on the canvas, or a grasshopper file with an sDNA_GH component on the canvas is first loaded, each component adds in Params for all its required Input and Output arguments (if the Params are not already present). These added Params include those of any extra automatically added tools if an auto_ option is true, that would other wise require separate components. Extra customisation can be carried out by adding in user specified Params too, that have the correct name of a supported option. Similarly any Params not being specified can be removed.

[note] The Config component tool always loads its options when placed or its Inputs are updated for any value of go. On the Unload_sDNA component, unload does the same thing as go.

Running individual tools.

Multiple sDNA_GH components can be chained together to run in sequence by connecting the OK Output Param of one component, to the go Input Param of the component(s) to be run afterwards. A Grasshopper Colour Gradient tool can be connected in between a Parse_Data component and Recolour_Objects component.

Component Execution Order.

Multiple sDNA_GH tools can be run from a single sDNA_GH component by setting any of the auto_options to true: auto_get_Geom, auto_read_User_Text, auto_write_Shp, auto_read_Shp and auto_plot_data, on a Config component, before placing the chosen sDNA_GH tool on the canvas.

Warning! If you did not create a config.toml file (in Installation step 12 above), and if you rely instead on Config components inside your .gh file itself to set option values, immediately before saving your .gh file, be sure to select the Config component determining any auto_ options, and press Ctrl + B (or from the pull-down menu select Edit -> Arrange -> Put To Back) to to send to the back, any components that should run first when you reload the .gh file. This ensures the Config component will run before other sDNA_GH components, which rely on settings controlled by it to configure themselves correctly.

Options.

sDNA_GH is highly customisable. This customisation is controlled by setting options. To give an option a value, connect a Grasshopper Param or text panel containing that value, to the Param with that option name on any sDNA GH component (except Unload sDNA). If a

Param is subsequently disconnected, its latest value will be remembered. Some Text Params can be cleared by connecting an empty Text Panel to them. Any option in a component can be read by adding an Output Param and renaming it to the name of the option. Similarly, any option in a component can be changed by adding an Input Param and renaming it to the name of the option, and connecting it to the new value. Entire options data structures (opts) may also be passed in from other sDNA_GH components as well, via normal Grasshopper connections.

Adding Component Input and Output Params.

To add a new Input or Output Param, zoom in on the component until plus and minus symbols can be seen between the params. Click on the plus symbol where you want the new Param. Right click the new Param's name (e.g. x, y or z for an Input Param, or a,b or c for an Output Param) to rename it to name of the desired option you wish to set.

Logging options

The logger is setup before Input Params are created on components, so to change the logging levels or to use a specific log file, log_file_level, log_console_level, working_folder, logs_dir or log_file must be set in an installation wide config.toml file, e.g. using a Config component. Supported values for logging levels are: DEBUG, INFO, WARNING, ERROR and CRITICAL.

Options override priority order

- 1. The component input Param options override options in a project specific options file (config).
- 2. A project specific options file (specified in config) overrides options from another sDNA_GH component (connected in opts).
- 3. Options from another sDNA_GH component override the installation-wide options file (e.g. %appdata%\Grasshopper\UserObjects\sDNA_GH\config.toml).
- 4. The installation-wide options file overrides the sDNA_GH hard-coded default options in main.py [^note] [note] Dev note: the options in main.py themselves override every individual tool's default options in tools.py.

Local meta options.

By default all sDNA_GH components share (and may change) the same global opts (module options, tool options, and *meta* options) in the main.py module. If only one of each tool is needed (and there is only one version of sDNA), that will suffice for most users.

Advanced users may give sDNA_GH components different options to the others, by de-

synchronising from the global options. De-syncing occurs if a component's *local meta options* (sync_to_module_opts, read_from_shared_global_opts) are false. *local metas* are like any other option, but shared using l_metas instead of opts, and not updated automatically from the main module options (as this would defeat their entire purpose).

Tools.

Common component input and output Params

OK This output is true when a component has executed successfully. **go** Set this input to true (e.g. from a boolean toggle component) to run a component's tool. **file** Specifies the path of a file to write to, or that was written to. **Data** Data must be a Data Tree 2 branches deep at the first level: a branch each for keys $\{0;0\}$ and values $\{0;1\}$. The two nodes of this structure should have a branch for each geometric object (so the nth's keys and values should have paths $\{0;0;n\}$ and $\{0;1;n\}$). The lists at these nodes must be of equal length. The mth key and value of the nth geometric object should be $\{0;0;n\}[m]$ and $\{0;1;n\}[m]$ respectively. Read_Shp supplies a Data Tree in this required format, if the data is read from User Text or from a Shapefile. Grasshopper's path tools can be used to adjust compatible Data Trees into this format.

Geom Accepts a list of geometric objects (Rhino or Grasshopper). Data trees of objects need to be flattened. gdm Accepts a Geometry-data-mapping, a python nested dictionary. The keys are the UUIDs of geometric objects. The values are also dictionaries, containing key/value pairs for use as User Text. opts Accepts an options data structure (a nested dictionary of named tuples) from another sDNA_GH component. Only of use if they are not synced to the global module options. config The path of a TOML file (e.g. config.toml) to be read in containing sDNA_GH options settings.

Support tools

Config (config)

To guarantee your components are setup correctly, this component must run before all others when your .gh file is loaded. To ensure this, select it and press Ctrl + B (or from the pull-down menu select Edit -> Arrange -> Put To Back) before saving.

Loads custom user options and configuration files (.toml). Saves options to a .toml file if go is true. If a .toml file is specified in save_to, it is saved to. Otherwise the default value of save_to is the installation-wide user options file. One is created if it does not already exist. This will overwrite existing files.

Read_Geom (get_Geom)

Reads in references to Rhino geometry (polylines) to provide them in the required form for subsequent sDNA_GH tools. Set the option selected to true, to only read the Rhino objects (of the specified type - polylines) that the user has selected. Similarly, specify layer = your_layer_name to only read Rhino objects from the layer named your_layer_name. To go back to selecting all layers, set layer to any value that is not the name of a layer.

The UUIDs of Rhino objects are converted to strings to preserve the references to them.

Shapefile tools

Write_Shp (write_shapefile)

Writes the DataTree in Data and list of objects (polylines) in Geom to a shapefile. If not specified in file, a default file name based on the Rhino doc or Grasshopper doc name is used (unless auto_update_Rhino_doc_path = false). overwrite_shp = true overwrites existing files; false or creates new files up to a maximum of max_new_files. WARNING! Shapefiles created with default names (due to no valid file path being specified in file by the user) will be deleted by subsequent sDNA tools if strict_no_del = false, overwrite_shp = false, and del_after_sDNA = true. To create a projection (.prj) file for the new shapefile, specify the path of an existing .prj file in prj. If no Data is supplied and auto_read_User_Text is true, this tool will first call read_User_Text. To work with sDNA, data records are only written to the Shapefile (associated with a shape corresponding to a Rhino / GH polyline) if its field matches the template string specified in input_key_str. The field name has a maximum of 10 characters long, and is taken from the {name} value (in the key name if it originated as User Text). To write all data with any key name (shorter then 11 characters) to the Shapefile, set input_key_str to {name}.

Read_Shp (read_shapefile)

Reads in polylines and associated data records from a shapefile. Creates new objects if new_geom = true or no objects corresponding to the shapefile are specified in Geom. Specify the path of the .shp file to read in file. WARNING! If a valid file path was not specified in file on a preceding Write_Shp component, and that file was used by an sDNA tool, Read_Shp deletes sDNA output shapefiles with default names if strict_no_del = false, overwrite_shp = false, and del_after_read = true. If a list of existing geometry is provided in Geom that corresponds to (is the same length as) the data records in the shapefile, only the data is read from the shapefile. Otherwise the shapes in the shapefile are outputted as new Grasshopper Geometry objects. The bounding box output bbox is provided to create a legend frame within Recolour_Objects (its value is calculated from the shape file). The abbreviations and field names from an sDNA results field file (if a file with the same name ending in .names.csv exists) are also read in, and supplied on abbrevs so that a drop-down list may be created, for easy selection of the data field for subsequent parsing and

plotting. If no separate Recolour_Objects Component is detected connected to the component's outputs downstream and auto_plot_data = true, Recolour_Objects is called afterwards.

Plotting tools

Parse_Data (parse_data)

Parse the data in a Data Tree of numerical data (in Data) from a specified field, for subsequent colouring and plotting. Be sure to supply the list of the data's associated geometric objects (in Geom), as legend tags and parsed values are appended to the output Data list and Geom list. Some classifiers sort the data into ascending order (if supplied, the geometry objects will then be reordered too, preserving their correspondence). To force a sort, according to field regardless, set sort_data to true. To make each parsed data point, take the same value as its class midpoint, set colour_as_class to true. Use this component separately from Recolour_Objects to calculate colours with a visible Grasshopper Colour Gradient component. Max and Min bounds can be overridden (in plot_max and plot_min). WARNING! Parsing is for the purpose of colourisation, e.g. in order to produce the desired result from Recolour_Objects. Therefore, although the inputted Data is not changed, the Data outputted almost certainly will be changed, so should be assumed to be false.

After parsing, the legend tags are the definitive reference for what each colour means, not the outputted data values. In particular, if colour_as_class = true, the parsed data will take far fewer distinct values than the number of polylines in a large network. To parse numerical data that uses a numerical format different to your system's normal setting (e.g. with a different radix character: ',' or '.' or thousands separator: ',' or '_'), set locale to the corresponding IETF RFC1766, ISO 3166 Alpha-2 code (e.g. fr, cn, pl).

Field to plot Specify the actual numeric data values to be parsed from all the provided 'User Text values' by setting field to the name of the corresponding 'User Text key'. Valid field values for sDNA output shapefiles are in fields.

Bounds The domain this data is parsed against can be customised by setting the options plot_min, plot_max, shifting it, widening it or narrowing it, e.g. to exclude erroneous outliers. If plot_min, plot_max are both numbers and plot_min < plot_max, their values will be used; otherwise the max and min are automatically calculated from the list of values in the 'User Text values' of Data corresponding to the 'User Text key' named in field. To go back to automatic calculation after an override, choose invalid values that satisfy plot_min >= plot_max. Set exclude to true to exclude data points lower than plot_min or higher than plot_max from the output altogether (and their corresponding objects from Geom). If exclude = false the plot_min, plot_max will be applied to limit the values of outlying data points (cap and collar).

Classes (bins / categories for the legend) Either, specify the number of classes desired in the legend in num_classes (the default is 7), or specify a list of the actual class boundaries desired in class_bounds manually. Note these are the inter-class bounds. Use plot_min for the lower bound of the bottom class and plot_max for the upper bound of the top class. There should be n-1 inter-class bounds, n classes and n+1 class bounds including the plot_max and plot_min.

If no valid inter-class boundaries are manually specified in class_bounds, sDNA_GH will automatically calculate them based on the following methods (each are valid values for class_spacing):

- quantile classify 'spikes' in the frequency distribution containing more data points than the normal class size, narrower than a specified width (in max_width). Then classify the remaining data values according to adjuster. Sorts the data.
- adjuster Sort the data and place inter class bounds in ascending order so that classes contain approximately the same number of data points, adjusting the inter-class boundaries to the closest gap, if one would otherwise be placed between identical data values.
- linear space the inter-class boundaries evenly between plot_min and plot_max.
- exponential space the inter-class boundaries between plot_min and plot_max but with a skewed spacing determined from an exponential curve (customisable base).
- log space the inter-class boundaries between plot_min and plot_max but with a skewed spacing determined from an logarithmic curve (customisable base).
- simple Uncomplicated quantile classification. Sort the data and divide it into classes containing approximately the same number of data points. Take no action if this places an interclass bound between identical values.
- max_deltas place the inter-class boundaries at the largest gaps between consecutive data points. Prone to distortion from outlying values. Sorts the data.

If after one of the above classification methods (especially simple), inter-class bounds have still been placed between indistinguishable data points (closer than tol), sDNA_GH can simply remove them (meaning there will be one few class for each) if remove_overlaps is set to true.

Legend class names Three customisable fields are provided in the options for the first, general and last legend tag names respectively: first_leg_tag_str = 'below {upper}', gen_leg_tag_str = '{lower} - {upper}', last_leg_tag_str = 'above {lower}'. A formatting string e.g. num_format = '{:.5n}' is applied to all numbers before displaying in the legend tags - it can be customised to set any desired number of decimal places or significant figures. If set, all must be valid Python format strings, with the supported named fields lower, mid_pt and upper, except num_format which supports a single unnamed field.

Re-normalisation Finally in order to produce a recolouring that has a set number of identical

colours (the same for each member of the same class) it is possible to assign the value of the midpoint of its class to each parsed data point. The parsed values may then additionally be renormalised, in order to tinker with the spread of colours against different colour gradients and other possible colourisations and applications, it is possible to 'renormalise' the parsed data points - the default value of re_normaliser is linear (for no re-normalisation) but exponential or log curves are also supported (with customisable base as above).

Finally, the errors raised if there are small classes or class overlaps can be suppressed by setting suppress_small_classes_error or suppress_class_overlap_error to true respectively.

Recolour_Objects (recolour_objects)

Recolour objects (and legend tags) based on pre-parsed and pre-normalised data, or already calculated colours (as RGB triples). If unparsed data is inputted, Parse_Data is first called. Custom colour curves are supported using a 3D quadratic spline between the triples of numbers: rgb_min, rgb_mid and rgb_max. Otherwise, use the Grasshopper Colour Gradient internally (via Node In Code) by setting Col_Grad to true and picking a setting from 0 to 7 for Col_Grad_num (0 : 'EarthlyBrown', 1 : 'Forest', 2 : 'GreyScale', 3 : 'Heat', 4 : 'Pink', 5 : 'Spectrum', 6 : 'Traffic', 7 : 'Zebra'). Set line_width to control the width of the line of Rhino geom objects (the default is 4).

Create a legend by connecting leg_cols, leg_tags and leg_frame to a Grasshopper Legend component. The coordinates of the corners of the Rectangle provided in leg_frame may be overridden by specifying leg_extent (xmin, ymin, xmax, ymax); alternatively any rectangle object can be passed into leg_frame on the GH Legend component itself. Custom legend tag templates and class boundaries are supported via four format strings (first_leg_tag_str, gen_leg_tag_str, last_leg_tag_str and num_format) as per Parse_Data.

To recolour Grasshopper geometry instead of Rhino Geometry (i.e. unbaked objects), connect the Data and Geom outputs to a Grasshopper Custom Preview component (line widths of GH objects cannot be increased).

User Text tools

Data tools

Read_User_Text (read_User_Text)

Reads all User Text from the list of Rhino objects in Geom. If auto_get_Geom = true, Read_From_Rhino is first called. If compute_vals = true, values starting and ending in % referring to a Rhino object's UUID (e.g. %<CurveLength("ac4669e5-53a6-4c2b-9080-

bbc67129d93e")>%) are computed using Rhino.RhinoApp.ParseTextField.

Write_User_Text (write_User_Text)

Writes User Text to Rhino objects, using a specified pattern for the keys. Specify the data tree to write in Data, and the list of Rhino objects to write to in Geom. The format of the User Text key can be customised in the Python format string output_key_str, accepting two named fields (e.g. = sDNA output={name} run time={datetime}).

A field specifying an originating Rhino object's UUID uuid_field will be omitted. If a key of that name already exists, it will be overwritten if overwrite_UserText is true. Otherwise a suffix will be appended to it, based on an integer counter and the format string in dupe_key_suffix, until a unique key name is found, up to a limit of max_new_keys (overwrite warnings can be suppressed by setting suppress_overwrite_warning to true).

sDNA Tools

Analysis tools

- sDNA tools run sDNA from the command line, using the Python interpreter in python.
- All sDNA tools try to load an sDNA installation. The first pair of sDNAUISPEc.py and runsdnacommand.py files matching the names in sDNAUISpec and runsdnacommand, found in a folder in sDNA_paths are loaded (if the corresponding sDNA is not already loaded). This is used to run the correct sDNA tool in the corresponding /bin sub folder, and to add Input Params to the sDNA component for each of its sDNA tool's inputs.
- By default an sDNA tool component will show all the possible inputs on its input Params. To show only the essential inputs instead (and make the components a lot smaller) set show_all = false.
- sDNA tools require a shapefile to be specified in file or input. If Write_User_Text is run beforehand and a file name is not specified, a default file name will be used.
- if the user Param make_advanced is true, all other unrecognised user params on the component will be added into the advanced config string.

If auto_write_Shp or auto_read_Shp are true, all sDNA components attempt to check, e.g. if any Write_User_Text or Read_User_Text components are already connected to its inputs (upstream) or outputs (downstream) respectively. If not, the sDNA component will run Write_User_Text or Read_User_Text before and after it. If all auto_ options are true, an sDNA component will take in geometry from Rhino directly, write it to a shapefile, run the analysis in sDNA, read in the output shapefile, and recolour the Rhino polylines.

WARNING! If a valid file path was not specified in file or input on a preceding Write_Shp component, and that file was used by an sDNA tool, sDNA components will delete input shapefiles with default names if strict_no_del = false, overwrite_shp = false, and del after sDNA = true.

The sDNA tool descriptions below are copied almost verbatim from the <u>sDNA manual</u>:

<u>sDNA Integral</u> (sDNAIntegral)

sDNA Integral is the core analysis tool of sDNA. It computes several flow, accessibility, severance and efficiency measures on networks.

This automatically calls other support tools, handling an entire Rhino geometry workflow from this one component, additionally running Read_Geom and Write_Shp before the sDNA tool itself, and then Read_Shp and Recolour_Objects afterwards (unless auto_write_Shp = false or auto_read_Shp = false respectively). WARNING! All sDNA tools will delete the shapefile named in input after it has been read in, if del_after_sDNA = true and strict_no_del = false (as they are by default).

To add or remove existing Geometry before the results file is read in (to control creation of new geometry objects), set auto_read_Shp = false and connect a Read_Shp component. To analyse a network of Grasshopper Geometry set auto_get_Geom and auto_read_User_Text to false. To access the data and geom objects before parsing and recolouring, auto_plot_data = false (and connect Parse_Data and Recolour_Objects components). This allows picking a results field from abbrevs to parse without repeating the whole analysis, and using a Grasshopper Colour Gradient component on the canvas to generate colours. Connect a Grasshopper Legend component to plot a legend. To recolour Grasshopper geometry instead of Rhino Geometry (i.e. unbaked objects), connect the Data and Geom outputs to a Grasshopper Custom Preview component.

To use sDNA's advanced config options in sDNA_GH, add in an input Param to an sDNA component with the same name as each advanced config option you wish to include (omitting a trailing equals sign and leaving the Param unconnected, unless you wish to provide a value for it); the advanced config string can then be saved to a config.toml file with an sDNA_GH Config component). Alternatively create an advanced config string manually. The sDNA tools in that component will gather all user-specified input Params and construct the advanced config string from them. Alternatively, prepare the and connect it to advanced. See the readme for the list of supported advanced config options.

Advanced config options for sDNA Integral

Option Default Description
startelev= Name of field to read start elevation from
endelev= Name of field to read end elevation from
metric= angular Metric – angular, euclidean, custom or one of the presets
radius= n List of radii separated by commas
startelev= Name of field to read start elevation from
endelev= Name of field to read end elevation from
origweight= Name of field to read origin weight from

destweight= Name of field to read destination weight from

origweightformula= Expression for origin weight (overrides origweight)

destweightformula= Expression for destination weight (overrides destweight)

weight= Name of field to read weight from. Applies weight field to both origins and destinations.

zonesums= Expressions to sum over zones (see zone sums below)

lenwt Specifies that weight field is per unit length

custommetric= Specified field name to read custom metric from

xytol= Manual override xy tolerance for fixing endpoint connectivity.

ztol= Manual override z tolerance for fixing endpoint connectivity.

outputgeodesics Output geometry of all pairwise geodesics in analysis (careful – this can create a lot of data)

output destinations Output geometry of all pairwise destinations in analysis (careful – this can create a lot of data). Useful in combination with origins for creating a map of distance/metric from a given origin.

outputhulls Output geometry of all convex hulls in analysis

outputnetradii Output geometry of all network radii in analysis

origins= Only compute selected origins (provide feature IDs as comma separated list). Useful in conjunction with outputgeodesicsm, outputdestinations, outputhulls, outputnetradii.

destinations= Only compute selected destinations (ditto)

nonetdata Don't output any network data (used in conjunction with geometry outputs)

pre= Prefix text of your choice to output column names

post= Postfix text of your choice to output column names

nobetweenness Don't calculate betweenness (saves a lot of time)

nojunctions Don't calculate junction measures (saves time)

nohull Don't calculate convex hull measures (saves time)

linkonly Only calculate individual link measures.

outputsums Output sum measures SAD, SCF etc as well as means MAD, MCF etc.

probroutes Output measures of problem routes – routes which exceed the length of the radius forcecontorigin Force origin link to be handled in continuous space, even in a discrete analysis. Prevents odd results on very long links.

nqpdn= 1 Custom numerator power for NQPD equation

nqpdd= 1 Custom denominator power for NQPD equation

skipzeroweightorigins Skips calculation of any output measures for origins with zero weight. Saves a lot of time if many such origins exist.

skipzeroweightdestinations 1 Zero weight destinations are skipped by default. Note this will exclude them from geometry outputs; if this is not desired behaviour then set skipzeroweightdestinations=0

skiporiginifzero= Specified field name. If this field is zero, the origin will be skipped. Allows full customization of skipping origins.

skipfraction= 1 Set to value n, skips calculation for (n-1)/n origins. Effectively the increment value when looping over origins.

skipmod= 0 Chooses which origins are calculated if skipfraction?1. Effectively the initial

value when looping over origins: every skipfractionth origin is computed starting with the skipmodth one.

nostrictnetworkcut Don't constrain geodesics to stay within radius. This will create a lot more 'problem routes'. Only alters behaviour of betweenness measures (not closeness).

probrouteaction= ignore Take special action for problem routes that exceed the radius by a factor greater than probroutethreshold. Can be set to ignore, discard or reroute. Reroute changes geodesic to shortest Euclidean path. Only alters betweenness output, not closeness. probroutethreshold= 1.2 Threshold over which probrouteaction is taken. Note this does not affect computation of probroutes measures, which report on all routes which exceed the radius length regardless of this setting.

outputdecomposableonly output only measures which are decomposable i.e. can be summed over different origins (useful for parallelization)

linkcentretype= Angular for angular analysis, Euclidean otherwise Override link centre types – angular or Euclidean

lineformula= Formula for line metric in hybrid analysis (see below)

juncformula= 0 Formula for junction turn metric in hybrid analysis (see below)

bidir Output betweenness for each direction separately

oneway= Specified field name to read one way data from (see note 1 below)

vertoneway= Specified field name to read vertical one way data from (see note 1 below)

oversample= 1 Number of times to run the analysis; results given are the mean of all runs.

Useful for sampling hybrid metrics with random components.

odmatrix Read OD matrix from input tables (a 2d table must be present)

zonedist= euc Set expression to determine how zone weights are distributed over links in each zone, or 0 to skip distribution (all lines receive entire zone weight)

intermediates= Set expression for intermediate link filter. Geodesics are discarded unless they pass through link where expression is nonzero.

disable= Set expression to switch off links (links switched off when expression evaluates nonzero)

outputskim Output skim matrix file

skimorigzone Origin zone field (must be text) for skim matrix

skimdestzone Destination zone field (must be text) for skim matrix

skimzone Skim matrix zone field for both origin and destination (sets both skimorigzone and skimdestzone)

bandedradii Divide radius into bands: for each radius only include links outside the previous radius

datatokeep= List of field names for data to copy to output

sDNA_Skim (sDNASkim)

Skim Matrix outputs a table of inter-zonal mean distance (as defined by whichever sDNA Metric is chosen), allowing high spatial resolution sDNA models of accessibility to be fed into existing zone-base transport models.

sDNA_Int_From_OD (sDNAIntegralFromOD)

A simplified version of sDNA Integral geared towards use of an external Origin Destination matrix. Note that several other tools (including Integral) allow Origin Destination matrix input as well.

The file must be formatted correctly, see Creating a zone table or matrix file. All geodesic and destination weights are replaced by values read from the matrix. The matrix is defined between sets of zones; polylines must contain text fields to indicate their zone.

sDNA Access Map (sDNAAccessibilityMap)

Outputs accessibility maps for specific origins, including metric between each origin-destination, Euclidean path length and absolute diversion (difference between Euclidean path length and crow flight path length, similar to circuity, notated here as 'Div').

The accessibility map tool also allows a list of origin polyline IDs to be supplied (separated by commas). Leave this parameter blank to output maps for all origins. If outputting "maps" for multiple origins, these will be output in the same feature class as overlapping polylines. It may be necessary to split the result by origin link ID in order to display results correctly.

sDNA Accessibility Map is a different interface applied to sDNA Integral, so will in some cases accept its advanced config options as well. To use sDNA's advanced config options in sDNA_GH, add in an input Param to an sDNA component with the same name as each advanced config option you wish to include (omitting a trailing equals sign and leaving the Param unconnected, unless you wish to provide a value for it). The sDNA tools in that component will gather all user-specified input Params and construct the advanced config string from them. Alternatively, prepare the advanced config string manually and connect it to advanced. See the readme for the list of supported advanced config options.

Preparation tools

<u>sDNA Prepare</u> (sDNAPrepare)

Prepares spatial networks for analysis by checking and optionally repairing various kinds of error. Note that the functions offered by sDNA prepare are only a small subset of those needed for preparing networks. A good understanding of Network Preparation is needed, and other (free) tools can complement sDNA Prepare.

The errors fixed by sDNA Prepare are:

- endpoint near misses (XY and Z tolerance specify how close a near miss)
- duplicate lines

- traffic islands (requires traffic island field set to 0 for no island and 1 for island). Traffic island lines are straightened; if doing so creates duplicate lines then these are removed.
- split links. Note that fixing split links is no longer necessary as of sDNA 3.0 so this is not done by default.
- isolated systems.

To use sDNA's advanced config options in sDNA_GH, add in an input Param to an sDNA component with the same name as each advanced config option you wish to include (omitting a trailing equals sign and leaving the Param unconnected, unless you wish to provide a value for it). The sDNA tools in that component will gather all user-specified input Params and construct the advanced config string from them. Alternatively, prepare the advanced config string manually and connect it to advanced. See the readme for the list of supported advanced config options.

Advanced config options for sDNA Prepare

Option Description startelev= Name of field to read start elevation from endelev= Name of field to read end elevation from island= Name of field to read traffic island information from. Anything other than zero will be treated as traffic island islandfieldstozero= Specifies additional data fields to set to zero when fixing traffic islands (used for e.g. origin or destination weights) data_unitlength= Specifies numeric data to be preserved by sDNA prepare (preserves values per unit length, averages when merging links) data_absolute= Specifies numeric data to be preserved by sDNA prepare (preserves absolute values, sums when merging links) data_text= Specifies text data to be preserved (merges if identical, concatenates with semicolon otherwise) xytol= Manual override xy tolerance for fixing endpoint connectivity ztol= Manual override z tolerance for fixing endpoint connectivity merge_if_identical= Specifies data fields which can only be merged if identical, i.e. split links will not be fixed if they differ (similar to 'dissolve' GIS operation)

<u>sDNA Line Measures</u> (sDNALineMeasures)

Individual Line Measures. Outputs connectivity, bearing, euclidean, angular and hybrid metrics for individual polylines. This tool can be useful for checking and debugging spatial networks. In particular, connectivity output can reveal geometry errors.

Geometric analysis tools

sDNA Geodesics (sDNAGeodesics)

Outputs the geodesics (shortest paths) used by Integral Analysis.

The geodesics tool also allows a list of origin and destination polyline IDs to be supplied (separated by commas). Leave the origin or destination parameter blank to output geodesics

for all origins or destinations. (Caution: this can produce a very large amount of data).

sDNA Geodesics is a different interface applied to sDNA Integral, so will in some cases accept its advanced config options as well. To use sDNA's advanced config options in sDNA_GH, add in an input Param to an sDNA component with the same name as each advanced config option you wish to include (omitting a trailing equals sign and leaving the Param unconnected, unless you wish to provide a value for it). The sDNA tools in that component will gather all user-specified input Params and construct the advanced config string from them. Alternatively, prepare the advanced config string manually and connect it to advanced. See the readme for the list of supported advanced config options.

SDNA_Hulls (sDNAHulls)

Outputs the convex hulls of network radii used in Integral Analysis.

The convex hulls tool also allows a list of origin polyline IDs to be supplied (separated by commas). Leave this parameter blank to output hulls for all origins.

sDNA Convex Hulls is a different interface applied to sDNA Integral, so will in some cases accept its advanced config options as well. To use sDNA's advanced config options in sDNA_GH, add in an input Param to an sDNA component with the same name as each advanced config option you wish to include (omitting a trailing equals sign and leaving the Param unconnected, unless you wish to provide a value for it). The sDNA tools in that component will gather all user-specified input Params and construct the advanced config string from them. Alternatively, prepare the advanced config string manually and connect it to advanced. See the readme for the list of supported advanced config options.

sDNA_Net_Radii (sDNANetRadii)

Outputs the network radii used in Integral Analysis.

The network radii tool also allows a list of origin polyline IDs to be supplied (separated by commas). Leave this parameter blank to output radii for all origins.

sDNA Network Radii is a different interface applied to sDNA Integral, so will in some cases accept its advanced config options as well. To use sDNA's advanced config options in sDNA_GH, add in an input Param to an sDNA component with the same name as each advanced config option you wish to include (omitting a trailing equals sign and leaving the Param unconnected, unless you wish to provide a value for it). The sDNA tools in that component will gather all user-specified input Params and construct the advanced config string from them. Alternatively, prepare the advanced config string manually and connect it to advanced. See the readme for the list of supported advanced config options.

Advanced config options for sDNA geometry tools

Calibration tools

sDNA_Learn (sDNALearn)

sDNA Learn selects the best model for predicting a target variable, then computes GEH and cross-validated R^2 . If an output model file is set, the best model is saved and can be applied to fresh data using sDNA Predict.

Available methods for finding models are (valid options for algorithm):

- Single best variable performs bivariate regression of target against all variables and picks single predictor with best cross-validated fit
- Multiple variables regularized multivariate lasso regression
- All variables regularized multivariate ridge regression (may not use all variables, but will usually use more than lasso regression)

Candidate predictor variables can either be entered as field names separated by commas, or alternatively as a *regular expression*. The latter follows Python regex syntax. A wildcard is expressed as .*, thus, Bt.* would test all Betweenness variables (which in abbreviated form begin with Bt) for correlation with the target.

Box-Cox transformations can be disabled, and the parameters for cross-validation can be changed.

Weighting lambda (weightlambda) weights data points by $y^{\lambda-1}$, where y is the target variable. Setting to 1 gives unweighted regression. Setting to around 0.7 can encourage selection of a model with better GEH statistic, when used with traffic count data. Setting to 0 is somewhat analogous to using a log link function to handle Poisson distributed residuals, while preserving the model structure as a linear sum of predictors. Depending on what you read, the literature can treat traffic count data as either normally or Poisson distributed, so something in between the two is probably safest.

Ridge and Lasso regression can cope with multicollinear predictor variables, as is common in spatial network models. The techniques can be interpreted as frequentist (adding a penalty term to prevent overfit); Bayesian (imposing a hyperprior on coefficient values); or a mild form of entropy maximization (that limits itself in the case of overspecified models). More generally it's a machine learning technique that is tuned using cross-validation. The r^2 values reported by learn are always cross-validated, giving a built-in test of effectiveness in making predictions.

Regularization Lambda allows manual input of the minimum and maximum values for regularization parameter λ in ridge and lasso regression. Enter two values separated by a

comma. If this field is left blank, the software attempts to guess a suitable range, but is not always correct. If you are familiar with the theory of regularized regression you may wish to inspect a plot of cross validated r^2 against λ to see what is going on. The data to do this is saved with the output model file (if specified), with extension .regcurve.csv.

<u>sDNA_Predict</u> (sDNAPredict)

Predict takes an output model file from sDNA Learn, and applies it to fresh data.

For example, suppose we wish to calibrate a traffic model, using measured traffic flows at a small number of points on the network:

- First run a Betweenness analysis at a number of radii using Integral Analysis.
- Use a GIS spatial join to join Betweenness variables (the output of Integral) to the measured traffic flows.
- Run Learn on the joined data to select the best variable for predicting flows (where measured).
- Run Predict on the output of Integral to estimate traffic flow for all unmeasured polylines.

Dev tool(s)

Unload_sDNA (Unload_sDNA)

Unload the sDNA_GH Python package and all sDNA modules, by removing them from Grasshopper Python's shared cache (sys.modules).

The next sDNA_GH component to run after this one (that's not also an Unload_sDNA) will then reload the sDNA_GH Python package and installation-wide options file (config.toml), and any specified options including a project specific config.toml, without otherwise having to restart Rhino to clear Grasshopper's cache. sDNA tools will also try to load an sDNA installation.

sDNA_General (sDNA_General)

Run any other component by feeding the name of it into the "tool" input param.

Self_test (selftest)

Runs the unit tests of the sDNA_GH module and launcher.py.

Not a tool in the same sense as the others (this has no tool function in sDNA). The name

Self_test (and variations to case and spacing) are recognised by the launcher code, not the main package tools factory. In a component named "Self_test", the launcher will cache it, then replace the normal RunScript method in a Grasshopper component class entirely, with a function (unit_tests_sDNA_GH.run_launcher_tests) that runs all the package's unit tests (using the Python unittest module). Unit tests of the functions in the launcher, can also be added to the launcher code.

License.

See <u>license.md</u>

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Developer manual.

Dependencies.

To bulk unblock files, to avoid unblocking every file manually it is necessary to install <u>Powershell</u>. Otherwise no additional dependencies are required. sDNA_GH is shipped with files from the following python packages included: <u>PyShp (MIT License)</u> "version: 2.2.0" <u>Toml (MIT License)</u> Latest commit 230f0c9 on 30 Oct 2020

Build instructions.

- 1. If sDNA_GH has not automatically found the sDNA installation you wish to build components for, place a Config component (the one with a lightbulb icon) and add its path to sDNA_paths.
- 2. Run build_components.bat (if necessary open it and adjust the paths to your local folders, and the paths in \dev\sDNA_build_components.gh).
- 3. For non-Github users, a good quality pdf of this file (README.md) can be created in VS Code with the extension: <u>print, PD Consulting VS Marketplace Link</u>. This will render the markdown file in your web browser. Print it to a pdf with the name README.pdf in the same directory (using Save to Pdf in Mozilla instead of Microsoft Print to Pdf will preserve the URLs in the links).
- 4. Manually create Unload_sDNA_GH and Readme.txt components if required.

- 5. Run create release sDNA GH zip.bat to create the zip file for release.
- 6. Note: The components are only GhPython launchers with different names and different docstrings. As much code as possible has been shifted into the python package and the other sDNA_GH Python package files. If no changes to the launcher code have been made and no new components/tools are required, a new release can simply reuse the .ghuser files from an old release, and the new release's zip files can be created simply by re running create_release_sDNA_GH_zip.bat.

To build new sDNA components.

sDNA_GH will attempt to automatically build components and user objects for the sDNA tools in an sDNAUISpec.py, that it doesn't already have .ghuser GH component / User Object file for. It will also look for an .png icon file with the same name as the tool Class in sDNAUISPec in \sDNA_GH\components\icons, and will parse this very file (README.md) for a tool description, to swap in as the launcher code's docstring (which will become the User Object and component descriptions, and its mouse over text). Therefore:

- for each new component: Add a description to this file, README.md starting on the line after (tool Class name) in brackets, ending in two blank lines to this very file. Save it to %appdata%\Grasshopper\UserObjects\sDNA_GH\README.md (overwriting the previous one).
- for each new component: Prepare an icon file and save it to %appdata%\Grasshopper \UserObjects\sDNA_GH\components\icons. 24x24 is recommended by the Grasshopper developers, but it seems fairly flexible see sDNA_Integral. A format compatible with .Net's System.Drawing.Bitmap Class is required. .png has been tested.
- Open a new Grasshopper canvas with sDNA_GH installed.
- Place an sDNA_GH Config component.
- Setup sDNA_GH to use the new version of sDNA by specifying it in sDNA_folders (following Installation step 12 above).
- Ensure make_new_comps is true.
- If necessary Recompute the sheet press F5.
- The new user objects for the new components will be automatically created, and added to the sDNA section of the Grasshopper Plug-ins Ribbon. Copy the relevant .ghuser file(s) from %appdata%\Grasshopper\UserObjects\, and paste them in \sDNA_GH\components\automatically_built in the repo. Place copies of the updated README.md file and new icon files in there too for posterity.

The supported data types for inputs (forced to lower case) are in sDNA_ToolWrapper.sDNA_types_to_Params in tools.py:

- fc = Param_FilePath
- ofc = Param FilePath

- bool = Param Boolean
- field = Param String
- text = Param_String
- multiinfile = Param FilePath
- infile = Param FilePath
- outfile = Param_FilePath

Misc

To compile C# code to a grasshopper assembly (.gha file): Install Visual Studio 2017 community edition with VB / C# / .Net workflow [https://developer.rhino3d.com/guides /grasshopper/installing-tools-windows/#fnref:3] Install Rhino & templates as above [https://developercommunity.visualstudio.com/t/net-framework-48-sdk-and-targeting-pack-in-visual/580235] Install .Net v4.8. Change .csproj target to v4.8 [https://stackoverflow.com/questions/58000123/visual-studio-cant-target-net-framework-4-8]

GHPython for .ghuser: Select GHPython component. Optionally compile to .ghpy. File -> Create User Object

[^0] The Python 2.7 download can be verified using this <u>certificate</u> and <u>Gpg4win</u>.

[^1] The entire source code for sDNA_GH is visible on Github. All the source code is also visible in the download itself as the component launcher and Python package is visible, except the .ghuser files which each contain the launcher code under a different name, and are compiled. It is a little repetitive, but see the Build Instructions above to build them for yourself from the source code.

[^2] This script is largely code from Ed Wilson of Microsoft's <u>Dev Blog</u> or try this <u>alternative</u> <u>method</u>)