Lecture 12:

Project: Reporting progress and (unexpected) challenges

COSC 526: Introduction to Data Mining Spring 2021



Project: Status



Project

Motivation Describe the motivation of your work. To build the motivation, you can answer these questions:

- What is the problem you are tackling?
- How is the problem solved today?

Contributions List between 2 and 4 contributions of your work. Contributions are bullet points that define your solution. E.g.,

- We build a system that
- We validate the system accuracy by
- We measure the performance of the system by ...
- Write a section of 150 200 words

Project

Tests List the type of tests (measurements) you will perform. E.g.,

- What are your metrics of success?
- Where do you run your tests?
- What tests do you perform?
- How many times do you run each test?
- What do you measure?
- Write a section of 250 350 words.

Milestones

- March 26: Define your project (DONE)
 - Describe the motivation of your work
 - List between 2 and 4 contributions of your work
 - List the type of tests (measurements) you will perform
- April 2: No lecture (DONE)

Milestones

- April 9: Create a new notebook with your solution (DONE)
 - Write down the steps of your solution in distinct text cells; add one or multiple cells (as needed) to hold your code for each step. You can leave these software cells empty for the moment. Expand the text cells describing your solution.
 - Add visualization cells that allow you to visualize results. You can leave these software cells empty for the moment.
 - Add software to the code cells that upload data from source and pre-process data.
 - Push your notebook into your GitHub repository as frequently as needed

Milestones (TODAY)

- April 16: Finalize software and complete the test run within your notebook
- April 23: Create your poster and get feedback, submit draft
- April 30: Submit your final notebook and poster in GitHub
- May 7: Submit your 2-page abstract in GitHub

Poster layouts

Poster 1

• Poster from:

http://betterposters.blogspot.com/2011/04/critique-breast-cancer-inhibition.html





O⁶-Benzylguanine Inhibits Tamoxifen Resistant Breast Cancer Cell Growth and Resensitizes Breast Cancer Cells to Anti-Estrogen Therapy

Joshua Smith', George C Bobustuc', Rafael Madero-Visbal', Jimmie Colon', Beth Isley', Jonathan Ticku', Kalkunte S. Srivenugopal and Santhi Konduri¹

Cancer Research Institute of M.D Anderson Cancer Center Orlando Texas Tech University Health Sciences Center, Amarillo, TX



Abstract

Endocrine therapies using anti-estrogens are least toxic and very effective for breast cancers, however, tumer resistance to monofera remains a sumbailing block for successful theraps, Based on our reent study on the involvement of the DNA requir protein MGMT in panerunic canner (Clin Cancer Res. 15, 6697, 2000), here, we investigated whether MGMT overexpression mediates tamosition resistance, Specifically, we determined whether administration of MGMT inhabitor (Orberapsquarine (RGI)) at a ron-twice dose above or in combination with the anti-estrogens (unmovider fiberetard) crutals human tamosifier mediate (Clin Control Co

MOAT expression was found to be increased in heast cancer cells relative to normal heast epithelial cells, Also, MOAT elsevies were significantly higher in tamostice resistant MCF-7 companies to the parter falls. Silicrain; of the E4s expression using a specific siRNA resulted in augmentation of MOAT mENA and protein levels by a fold. We also observed an inverse correlation between MOAT and part in invest cancer cell files. Increase consequent of the contract contract of the most contract cells for the contract contract of the contract c

Introduction

Recent advances in breast cancer research have identified key pathways involved in the repair of DNA damage induced by chemotherapeutic agents. The ability of cancer cells to recognize DNA damage and initiate DNA repair is an important mechanism for therapeutic resistance and has a negative impact on therapeutic efficacy. A number of DNA-damaging alkylating arents attack the nucleonbilic O* position on quanine, forming mutagenic and highly cytotoxic interstrand DNA crosslinks. The DNA repair enzyme O6-alkylguanine DNA alkyltransferase (AGT), encoded by the gene MGMT, repairs alkylation at this site and is responsible for protecting both tumor and normal cells from alkylating agents. MGMT is expressed constitutively in normal cells and tissues. In breast tumors, MGMT gene expression is elevated and levels are up to 4-fold higher than in the normal breast. Interestingly, it has been shown that tamoxifen accelerates proteasomal degradation of MGMT in human cancer cells. In 1991, Pegg, Moschel, and Dolan observed that Obenzylguanine (BG) inhibited AGT and potentiated the cytotoxicity of both chloroethylating agents and methylating agents. In a series of important observations, they fully characterized the interaction between BG and AGT and its therapeutic impact. They showed that BG binds AGT, transferring the benzyl mojety to the active-site cysteine [29]. The reaction is very rapid and more potent than any other previously known AGT inhibitor. BG is not incorporated into DNA in living cells and reacts directly with both cytoplasmic and nuclear AGT. Because BG is a psuedosubstrate for MGMT which results in the covalent transfer of benzyl group to the active site cysteine, the MGMT protein is degraded after each reaction. This stoichiometric reaction mechanism effectively depletes the AGT content in tumors and the associated repair of alkylation damage. BG is currently undergoing clinical trials in various cancers to increase the efficacy of alkylating agents

Interestingle, several observations suggest an inverse correlation between the levels of MOMT and grg tumor suppressed perties where widely per gar suppresses transcription of human MOMT expression. Informatively, per function is often inactivated or suppressed in human cancers; therefore, restention of set-gar, activity is essential for the success of some texturents. However, whether or not this is meditated by suppression of MOMT expression has yet to be determined. To date, the cross-talk between MOMT and ER-alpha (and the link to pgc spression) has not been explored in drug (i.e., tumofilar) resistant breast tumors. The anti-stronget nameline is the most commonly used treatment for patients with estrogen receptor positive breast cancer. Although many patients benefit from tanonien in the adjuvant and meditative scripting receptor positive breast cancer. Although many patients benefit from tanonien in the adjuvant and meditative visua to investigate the mechanisms of anti-entropen drug resistance and to design new therapeutic strategies for the interesting patients. The situation of the control of the patient of the control of the patient of the control of the patient of the patient

Results

Prolonged Treatment of Tamoxifen Increases MGMT Expression: We developed a tamoxifen resistant MCP2cell line by using prolonged treatment of tamoxifen on the parental ER-positive breast cancer cell line, MCP2-7, Tamoxifen-resistant MCP2-7 cells proliferate at rates similar to the parental MCP2-7, Prolonged treatment of tamoxifen onto MCP2-7 cells for gold (Fig. 1).

Knocking Down ERE Enhances MGMT Expression in Tamoxifen Resistant Breast Canner Cells It is not known whether ERe and MGMT transcriptionally regulate each other in tamoxifen resistant breast cancer cells. We therefore investigated by the contraction of the con

Transcriptional Regulation Between MGMT and pgg; Previous), it was reported that pgg negatively regulates MGMT in breast cancer cells. Therefore, we addressed whether or not silencing the pgg enhances endogenous MGMT transcription. Temosities at MCT-0 with wester transferred with either pgg allies, Mgg;320 (Fig.2C) or 100 km and 100 km and



With In Figure 1. MCF-7 parental and tamosifien resistant MCF-7 cell peliets were prepared, proteins were insulated and MGMT expression was detected by western blost analysis. Tamosifier presistant MCF-9 tamoster cells significantly increased MGMT expression compared to MCF-9 parental

O'-Benzojanaine Flays a Dual Role in Tamosifon Resistant MCF- Cella: Contrasting with the experiments above, next, we studied whether on on thocking down MGMT and any effect on ERe transcription. As expected, Innoving down MGMT decreased MGMT gene transcripts. However, it was interesting to find that EEs gene transcription uses also reduced after MGMT effecting. (Fig. 23). These district formatter than 100 miles and builty of the Cell and the Section of the Cell and th

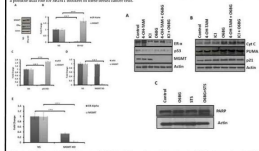


Figure 8, 10.7 Transide ministra MTG+ cells were transidened with Elea 6500, minorshi (Elea MTG) and Sallek (annuelle MTG), and elle were have resulted thy not received to the state of th

Figure 2, 4.0 Temestire moistant MCPe brosst entere cells were treated in presente an absence of EG (e.g. appl and his post treatment 4 oFLTMM (caM), ESZ (p,M) either above or in combination with ISC. 2ab post treatment cells were harvested and predicts were inducted most where his straight was performed. (A) ESZn pp3, and MOMT expensions (B) Cytechrone C, PUSA, and pa1 was determined by weaterm bot anaboles (C) transifient evisions MCPy cells were transied with or without RG in appl and later treated with stanosportin (5 pd/L) for 6 hr PAPE Greenage was determined by western blot anaboles.

Od-Benzylguanine Modulates pg3 Down-Stream Targeted Protein Expressions: Encouraged by the results reported, we are subjected for effect of combination therapy on endogenous MGMT, pg3, and Eng protein expressions. See speeds, 160 Celevated MGMT expression, while combination therapy (e.01-TAM or ICI combined with BG) significantly decreased both MGMT and Elke expressions, while does not combination with tumorision red Cel decreased Elke expression, where stamoufed nation and ICI continues are suppressionally the protein service of the expression in the protein significant protein

OG-Benzylgannine Modulated Transcriptional Targets in Tamouffen Resistant Breast Cancer Colls: The effect of combination therapy on endogenous MOHT GRNA levels that anti-settogens (TAM/ICI) increased the MOST expression while the combination therapy decreased it generally a supersion while the combination therapy decreased it compared to control with all these treatments (Fig.4a). Surprisingly, p21 and FUMA mRNA was significantly increased in the presence of combination tententist (Fig.4a) transcription was affected by the drug combinations in breast cancer cells (Fig. 2a three).

O6-Benzylguanine Enhances pzi Transcriptional Activity in Tamscriptional Activity in Tamscriptional Resistant Breast Cancer Cells In order to investigate the effect of 86 on pcg Junction, we performed hederiene reporter assays. Tamscrifen resistant promoter construct in presence or absence of 80 (target gene pcg.). These results clearly demonstrate that BG significantly enhanced pzi transcriptional activity by 4-5 fold in these cells (Fig. 40).

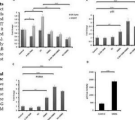
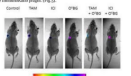
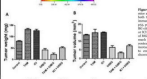


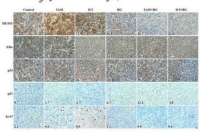
Figure a, Tancadra resistant MCT- front causer cells were treated in pressure or shown or fife (up ping the dath or 40 Hannodien and XI (pM)) was often after ori combinations with Kin of any fit or the other cells caused by the cells were harvested and table BNA was feelent, 6.0 MGMT and BRs (B) pin transcription. OF TMX transcription was doctomized by grift PCRC 4.0 Hannodien and KI falcone MGMT transcription. Bindood PVMA and pin transcription. BND immodien resistant MCT- breast cancer cells were transfered with pulse, ememorate and 60 later tracted with Euro and after cells were harvested, part transcription.

Go-Benylguanine Ishibits Tamosifon Resistant Breast Cancer Cell Growth and Increase Resistant Remast Cancer Cell Sensitistiy to Ant-Habstragon Therapy (TAM)/CI). Petalde necropgo revealed that all the mice had tumors in the breast. The data summarised in Table 1 show the duily BG alone or in combination with their weedly tamonised present mediant tumor volume and weight as compared with that seen in tamosifon/ICI treated and control mice. The combination of BG with tamosifien or ICI produced that seen in tamosifien/ICI treated and control mice (8, span 4, span 4, span 5, span 4, span

Histology and HC Analysis: We next determined the in view effects of BC (alone or in combination) with monofier/HC. Tumous havesed from different tentament groups were processed for routine histological and HC analysis. Tumous from mice treated with BC daine or in combination with tumoulies/HC exhibited a significant processed of the second second second second second second second second second expression was no much altered in these treatment groups. In sharp contrast, the expression of grant significantly increased in tumors from mice treated with BC either above or in combination with tumoulies/HC. He immunication gripping, Fig. 20.







- Conclusions

 1. In the present study, we observed that prolonged treatment with anti-estrogens causes drug resistance by inducing the DNA repair protein O*-metholyaumine DNA methottransferases (MGMT).
- Decreasing the expression of MGMT by exposing breast cancer cells to BG sensitined these cells to anti-extragen therapy (tamoxifen and ICI 182,786).
 We also observed that combination therapy of anti-estrogens and MGMT blockers not only overcame the MGMT derived drug (tamoxifen and ICI) resistance but also increased the efficacy of anti-estrogen therapy by decreasing estrogen receptor expression and restoration of the functional activity of per in amoxifering.

resistant breast cancer cells.

4. Combination therapy inhibited tamoxifen resistant breast tumor growth in vivo.

Acknowledgements

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O⁶-Benzylguanine Inhibits Tamoxifen Resistant Breast Cancer Cell Growth and Resensitizes Breast Cancer Cells to Anti-Estrogen Therapy

Joshua Smith', George C Bobustuc', Rafael Madero-Visbal', Jimmie Colon', Beth Isley', Jonathan Ticku', Kalkunte S. Srivenugopal and Santhi Konduri¹

*Cancer Research Institute of M.D Anderson Cancer Center Orlando *Texas Tech University Health Sciences Center, Amarillo, TX



Abstract

Endocrine therapies using anti-estrogens are least toxic and very effective for breast cancers, however, tumor resistance to amoxifen remains a stumbling block for successful therapy. Based on our recent study on the involvement of the DNA repair protein MGMT in pancreatic cancer (Clin Cancer Res. 15, 6087, 2000), here, we investigated whether MGMT overexpre nediates tamoxifen r or 10%-beneylguanine (BG)? at a non-toxic disils human tamoxifen resistant breast cance

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need abstracts the ER-u expression was accompanied by poxifen or fulvestrant decreased ER-u expression, whereas tamoxifen alone and fulvestrant alone increased and decreased the same respectively

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However, all these treatments increased the partie mRNA and protein expression significantly. BG inhibited tamoxifes resistant breast cancer growth in a dose-dependent manner and it also resensitized resistant breast cancer cells to antiestrogen therapy (TAM/ET). These combinations also enhanced the cytochrome C release and the PARP cleavage, indicative of aportosis. In breast cancer renografts, BG alone or a combination of BG with tamoxifen or fulvestrant caused significant tumor growth delay and immunohistochemistry revealed that BG inhibited the expression of MGMT, ER- u, ki-67 and nervased pay's staining. These findings suggest that MGMT inhibition may provide a novel and effective approach for overcoming tamoxifen resistance

Introduction

Recent advances in breast cancer research have identified key pathways involved in the repair of DNA damage induced by chemotherapeutic agents. The ability of cancer cells to recognize DNA damage and initiate DNA repair is an important mechanism for therapeutic resistance and has a negative impact on therapeutic efficacy. A number of DNA-damagin alkylating agents attack the nucleophilic O^o position on guanine, forming mutagenic and highly sytotoxic interstrand DNA rosolinks. The DNA repair enzyme O'-alkylguanine DNA alkyltransferase (AGT), encoded by the gene MGMT, repairs alkylation at this site and is responsible for protecting both tumor and normal cells from alkylating agents. MGMT is and levels are up

to 4-fold higher than inhibited AGT and po showed that BG binds more potent than any transfer of benzyl grou reaction mechanism e

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inhibition of MGMT by BG significantly improves TAM-sensitivity.

inactivated or suppress the success of some treatments. However, whether or not this is mediated by suppression of MGMT expression has yet to be determined. To date, the cross-talk between MGMT and ER-alpha (and the link to p53 expression) has not been explored in drug (i.e., tamoxifen) resistant breast tumors. The anti-estrogen tamoxifen is the most commonly used treatment for patients with estrogen receptor positive breast cancer. Although many patients benefit from tamoxifen in the adjuvant and metastatic settings, resistance to this endocrine therapeutic agent is an important clinical problem. The primary goal of present study was to investigate the mechanisms of anti-estrogen drug resistance and to design new therapeutic strategies for ng this resistance. The results show that MGMT expression is increased in TAM-resistant breast cancers and

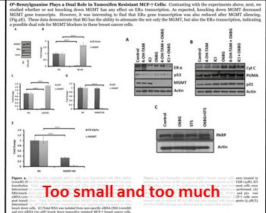
Prolonged Treatment of Tamoxifen Increases MGMT Expression: We developed a tamoxifen resistant MCF-7 cell line by using prolonged treatment of tamoxifen on the parental ER-positive breast cancer cell line, MCF-7 Tamoxifen-resistant MCF-7 cells proliferate at rates similar to the parental MCF-7. Prolonged treatment of tamoxifer onto MCF-7 cells increased MGMT expression compared to parental MCF-7 cells by 2 fold (Fig.1).

Knocking Down ERa Enhances MGMT Expression in Tamoxifen Resistant Breast Cancer Cells: It is not known whether Ellis and MCMT resourcistionally regulate each other in tamoxifen resistant breast cancer cells. We therefore investigated thether down regulation of EEa has any effect on endogenous MGMT expression in these cells. As expected, downregulation of ERB using specific siRNA significantly reduced ERB protein levels in these cells. Western blot analysis was performed and the results in the left panel (Fig. 2A) shows that silencing of ERs increases MGMT expression in these cells, and interestingly, the results in the right panel (Fig.2B) show increased MGMT mRNA levels were increased as assessed by qRT-PCR. These data suggest that ERO-mediated signaling functions to repress MGMT gene expression in

Transcriptional Regulation Between MGMT and p53: Previously, it was reported that p53 negatively regulates MGMT in breast cancer cells. Therefore, we addressed whether or not silencing the p53 enhances endogenous MGMT transcription. Tamoxifen resistant MCF-7 cells were transfected with either p53 siRNA (p53-KD) (Fig.2C) or MGMT siRNA (MGMT-KD) (Fig.2D) along with Non-specific siRNA (NS). MGMT spression was consistently increased in p53 knock down cells, with different speriments showing a - fold augmentation (Fig. 2A) and as expected, knocking down MGMT decreased MGMT transcripton where as pg3 mRNA levels were unaffected in Figure 1. MCF-7 present and tensile MGMT become cells (Fig. a-D). These results confirm that next can results MGMT as mistant MCF-7 of polisis were properly MGMT knockdown cells (Fig. 2D). These results confirm that p53 can regulate MGMT at



connectes was detected by western blot nion command to MCE/7 nan

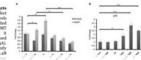


06-Benzylguanine Modulates p53 Down-Stream Targeted Protein Expressions: Encouraged by the results reported, investigated the effect of combination therapy on endogenous MGMT, p.g., and ERD protein expressions. As expected, BG decreased
MGMT expression, while combination therapy (4-OH-TAM or ICI combined with BG) significantly decreased both MGMT and ERs pressions. BG alone or in combination with tamoxifen or ICI decreased ER-0 expression, whereas tamoxifen alone and ICI alone reased and decreased the same respectively (Fig.3A), pg3 expression was slightly altered after RT treatment. The reduction in pg; pression by ICI alone was reversed when BG was combined (Fig. 3A). We investigated the effect of BG on proteins which are involved in cell cycle regulation, apoptosis in tamoxifen resistant breast cancer cells. All these treatments significantly increased the pgr protein expression (Fig.3B). PUMA expression was also increased with these treatments. Hence, PUMA may have translocated to the sitochondria, cytochrome C is released (Fig.3B), and apoptosis was triggered in these cells in presence of combination therapy. FARF deavage is seen in BG treated cells in presence of staurosporin as an indicative of apoptosis (Fig.yC). Therefore, this data suggest that BG promotes cell cycle arrest and can induce apoptosis by modulating pgg function.

06-Benzylguanine Modulated Transcriptional Targets in Tamoxifen Resistant Breast Cancer Cells: The effect of combination therapy on endogenous MGMT mRNA levels ws aso stuid. Quantitative real-time PCR (qRT-PCR) resulted that anti-estrogens (TAM/ICI) increased the MGMT expression while the combination therapy decreased it compared to control levels. ERg transcription was decreased compared to controls with all these treatments (Fig.4A). Surprisingly, not and PUMA mRNA was significantly increased in the presence of combination treatments (Fig. 4)

SDMT and pijj itunoription was determined by qRT-PCE. (3) Total RNA was related from mon-specific siRNA (30) (100mM) and MGMT siRNA (100mM) knock

town turnsiles resistant MCE: * broast currer ords. MCMT and not true



Caption not aligned with figure

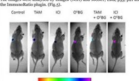
In order to investigate the effect of BG on p53 function performed luciferase reporter assays. Tamoxifen i MCF-7 breast cancer cells were transfected with promoter construct in presence or absence of BG (targe one of p53). These results clearly demonstrate the BG significantly enhanced not transcriptional artivity by in these cells (Fig.4D).





48h and later a CRI tamoulles and KT (saM) was either along or in combination with BC and sub la ordic work harvested and total RNA was included, OU MGMT and ERn (III) are transcription (IC) PGT transcription was determined by qRT PCR. a OR temosiles and RT induces MGMT transcription. induced PCSAA and pits transcription. (2) Tamoulles resistant MCF+2 broast nancer cells were transfer with part-for constitut and 6th later-treated with BG and agh later-cells were harvested, pay transcription activity was resistantly interested in SG in these cells. cylguanine Inhibits Tamoxifen Resistant Breast Cancer Cell Growth and Increase Resistan ncer Cell Sensitivity to Anti-Estrogen Therapy (TAM/ICI): Detailed necropsy revealed that a tumors in the breast. The data summarized in Table 1 show the daily BG alone or in combination by tamoxifen/ICI significantly decreased median tumor volume and weight as compared with that I treated and control mice. The combination of BG with tamoxifen or ICI produced the an tumor volume as compared with control mice (83.99 mm), 9.33 mm3 (TAM+BG) \$3.99 mm³, 31.60 mm³ (ICI+BG), respectively; p<0.0001). Tumor weight was also greated with combination therapy as compared with control mice (81.23 mg, 22.30 ing (ICI+BG), respectively, p-00.0005), (Table.1). Body spared with control mice. No visible liver metasta scope) in all treatment groups.

in vivo effects of BG (alone or in combination) wit roups were processed for routine histological and IHC malysis. Tumors from mice treated with BG alone or in combination with tamoxifen/ICI exhibited a significant fecrease in MGMT, ERO, ki-67 as compared with tumors treated with tamoxifen/BCI alone or control group. pgg repression was not much altered in these treatment groups. In sharp contrast, the expression of pax was significantly increased in tumors from mice treated with BG either alone or in combination with tamoulien/ICI The images were analyzed by ImageJ (NIH) and MGMT, ERG, pg3, p21 and ki-67 expressions were quantified by



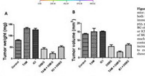
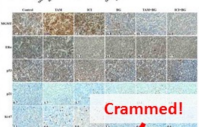


Figure 5. Tumors were harvasted from control nior and mice treated with temesitins/VL BG, no both temesitins/RC and BG. The nections were immunositation for expression of MOMT, Elst, pG, pm and 16-49. Temesor from nior to mosted with BG offlow alone or in constitution with termsoline of MCMT. Title and little. ptp expension was no such altered in these treatment groups, In shar

cells to BG sensitized these cells to anti-



Conclusions 1. In the present study, we observed that prolonged treatment wit inducing the DNA repair protein Of-methylguanine DNA methy resference (MGMT)

estrogen therapy (tamoxifen and ICI 182,780). We also observed that combination therapy of anti-estrog and MGMT blockers not only overcame the increased the efficacy of anti-estrogen therapy of the functional activity of pgg in tamoxifen-MGMT derived drug (tamoxifen and ICI) resistance but by decreasing estrogen receptor expression and reston

Acknowledgements

resistant breast cancer cells. 4. Combination therapy inhibited tamoxifen resistant breast tumor growth in vivo

2. Decreasing the expression of MGMT by exposing breast can-

More about how to prepare posters

• The following set of slides are from this extended lecture:

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https://urc.ucdavis.edu/sites/g/files/dgvnsk3561/files/local_resources/documents/p df documents/How To Make an Effective Poster2.pdf
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Project progress

- Jerome Kovoor and Shree Neupane: Analyzing the effect of climate change on global food production using K-means clustering or DBSCAN
- Fabian Fallas Moya: How many are good enough?
 Finding the best number of annotated images into a self-training algorithm
- Gerald Jones: Subgroups and Factors of ESRD



- Xinlan Jia & Candice Chen: US Airbnb Price Prediction
- Zhixiu Lu: Using Microbiomes to Predict Environmental Factors via Machine Learning Approaches
- Azarang Asadi: Motor control quantification using lower-limb body kinematics
- Tommy Li: Applications of Variational Autoencoders in Analyzing Ferroelectric Domains



- Georgia Channing: Traces in the Noise: Identifying Invalid
 Vessel Paths
- Anuj Gautam: Add a symbol and try again: A comprehensive study of password policies
- Mirka Mandich & Jake Maeker: Machine Learning Applied to HIT-SI Spheromak Data
- Carter White: Impact of Champion Selection on League of Legends Rank



- Levente Dojcsak: Predicting the Development of CKD and Identifying Preventative Treatments
- Konstantinos Georgiou: Analyzing and predicting bottlenecks on the distribution of COVID-19 Vaccines
- Michael Wermert: Stock Predictor: Using Machine Learning to Predict Stock Market Behavior

